Contemporary H3N2 influenza viruses have a glycosylation site that alters binding of antibodies elicited by egg-adapted vaccine strains

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H3N2 viruses continuously acquire mutations in the hemagglutinin (HA) glycoprotein that abrogate binding of human antibodies. During the 2014–2015 influenza season, clade 3C.2a H3N2 viruses possessing a new predicted glycosylation site in antigenic site B of HA emerged, and these viruses remain prevalent today. The 2016–2017 seasonal influenza vaccine was updated to include a clade 3C.2a H3N2 strain; however, the egg-adapted version of this viral strain lacks the new putative glycosylation site. Here, we biochemically demonstrate that the HA antigenic site B of circulating clade 3C.2a viruses is glycosylated. We show that antibodies elicited in ferrets and humans exposed to the egg-adapted 2016–2017 H3N2 vaccine strain poorly neutralize a glycosylated clade 3C.2a H3N2 virus. Importantly, antibodies elicited in ferrets infected with the current circulating H3N2 viral strain (that possesses the glycosylation site) and humans vaccinated with baculovirus-expressed H3 antigens (that possess the glycosylation site motif) were able to efficiently recognize a glycosylated clade 3C.2a H3N2 virus. We propose that differences in glycosylation between H3N2 egg-adapted vaccines and circulating strains likely contributed to reduced vaccine effectiveness during the 2016–2017 influenza season. Furthermore, our data suggest that influenza virus antigens prepared via systems not reliant on egg adaptations are more likely to elicit protective antibody responses that are not affected by glycosylation of antigenic site B of H3N2 HA.

Influenza vaccines possess antigens from one H1N1 strain, one H3N2 strain, and one or two influenza B strains. H3N2 viruses began circulating in humans in 1968. Most neutralizing antibodies (Abs) recognize antigenic sites on the globular head of HA (designated sites A–E), while rare Abs bind to the more conserved HA stalk or sialic acid binding domains (5). Most H3N2 vaccine mismatches from 1968 to 2013 have been attributed to mutations in antigenic site B of HA (6). None of the site B mutations that emerged during this time period have led to new glycosylation sites on HA (6, 7). This is surprising, given that the addition of glycans on HA can dramatically affect Ab binding (8–13). Instead, new glycosylation sites have repeatedly emerged and fixed in other antigenically important regions of H3 (14).

Vaccine effectiveness was extremely low during the 2014–2015 influenza season (15). During that season, influenza vaccines possessed antigens from a 2012 H3N2 virus that belonged to the 3C.1 HA genetic clade, while the majority of circulating H3N2 strains belonged to the 3C.2a and 3C.3a genetic clades (16). Both the 3C.2a and 3C.3a viruses differed at residues in HA antigenic site B compared with the 2014–2015 H3N2 vaccine strain (17). Notably, 3C.2a viruses that circulated during the 2014–2015 season possessed a new predicted glycosylation site in HA antigenic site B (17), and the 2014–2015 influenza vaccine exhibited especially low effectiveness against this clade (18). In an effort to avoid an antigenic mismatch, the 2016–2017 influenza vaccine was updated to contain antigens from a 3C.2a H3N2 virus isolated in 2014 (19). The majority of H3N2 viruses that circulated in the Northern Hemisphere during the 2016–2017 influenza season were 3C.2a viruses; however, an interim estimate of vaccine effectiveness was only 43% against medically attended H3N2 infections (20). Vaccine effectiveness was especially low in individuals that were 18–49 y old (20). It is unclear why there was variable vaccine effectiveness during the 2016–2017 influenza season, given that the vaccine strain appeared to be well matched to most circulating strains. The majority of antigens for influenza vaccines are prepared in fertilized chicken eggs, and the 2016–2017 egg-adapted 3C.2a vaccine strain lacks the site B glycosylation site that is present on circulating 3C.2a H3N2 strains (21). Here, we completed a series...
of studies to determine whether the difference in glycosylation of HA antigenic site B of H3N2 vaccine strains and circulating strains contributed to a previously unrecognized vaccine mismatch during the 2016–2017 influenza season.

Results

Recent H3N2 Viruses Possess a Glycosylation Site in Antigenic Site B of HA. A K160T HA mutation rapidly rose to fixation during the 2014–2015 influenza season, and nearly all currently circulating H3N2 viruses possess threonine (T) at HA residue 160 (Fig. 1A). The K160T HA mutation is predicted to introduce an N-linked glycosylation site in antigenic site B of HA (Fig. 1B and Table S1). H3N2 viruses with the K160T HA mutation grow poorly in chicken eggs (21), and the 2016–2017 egg-adapted H3N2 vaccine strain possesses a T160K HA reversion mutation [reference: Global Initiative on Sharing All Influenza Data (GISAID) isolate ID EPI_ISL_189811]. We used reverse genetics to create H3N2 viruses possessing HAs with T160 and K160, and we completed Western blot analyses to determine whether the T160 and K160 HAs migrate differently in SDS-PAGE gels. HAs with T160 migrated with a higher molecular weight compared with HAs with K160 (Fig. 1C, Left). The HAs migrated similarly after PNGase treatment (Fig. 1C, Right), indicating that the difference in HA mobility in the absence of PNGase is due to differences in N-linked glycosylation.

To determine whether the addition of this new antigenic site B glycan in HA affects antigenicity, we tested the binding of a panel of 26 human monoclonal antibodies that were elicited against an H3 virus from 2009, which lacks the new glycosylation site in antigenic site B of HA. All of these monoclonal antibodies bound to the HA of a 2009 virus (Fig. 2). The majority of these monoclonal antibodies (77%) bound to a 2014 virus with the egg-adaptive K160 HA (which lacks the glycosylation site), while only a few (23%) bound to a 2014 virus with T160 (which possesses the glycosylation site) (Fig. 2B). Raw binding data of a monoclonal antibody (024-10128-3C04) that binds efficiently to virus with K160 HA but not T160 HA is shown in Fig. 2B. As a control, we also measured binding of an HA stalk-reactive monoclonal antibody that efficiently recognizes virus with either T160 HA or K160 HA (Fig. 1C). Together, these data demonstrate that current circulating H3N2 influenza viruses possess a new glycosylation site in antigenic site B of HA that affects antigenicity, and that this glycosylation site is not present in current egg-adapted H3N2 vaccine strains.

Antibodies Elicited in Ferrets Exposed to the 2016–2017 Egg-Adapted H3N2 Vaccine Strain Poorly Neutralize a Circulating H3N2 Viral Strain.

We next completed a series of experiments to determine whether the current egg-adapted H3N2 vaccine strain (that possesses K160) elicits different types of antibodies compared with the current circulating H3N2 viral strain (that possesses T160). We infected ferrets with viruses possessing either K160 HA or T160 HA, and we completed foci reduction neutralization tests (FRNTs) using sera collected from these animals 28 d after infection. We completed FRNT assays rather than conventional hemagglutination–inhibition assays since 3C2.a H3N2 viruses inefficiently agglutinate red blood cells. Antibodies elicited by the egg-adapted vaccine strain possessing K160 HA recognized the egg-adapted vaccine strain fourfold to eightfold more efficiently than the currently circulating H3N2 strain that possesses T160 HA (Fig. 3A). This indicates that a large proportion of ferret antibodies elicited by the current egg-adapted H3N2 vaccine strain recognize the unglycosylated antigenic site B of HA. Interestingly, antibodies elicited by infection with the current circulating H3N2 viral strain possessing T160 HA recognized viruses with T160 HA and K160 HA equally (Fig. 3B). This is consistent with the hypothesis that the new HA glycosylation site effectively “shields” antigenic site B of HA, and that antibodies elicited by the viral strain possessing the new HA glycosylation site recognize epitopes not involving antigenic site B of HA.
Some individuals in our study mounted strong antibody responses against T160 HA following vaccination (Fig. 41). Antibody increases against the K160 HA were similar following vaccination with Fluzone and Flucelvax (Fig. 4B). These differential effects of the vaccines remained statistically significant after adjusting for age, prevaccination titer, and vaccination history (Table S3). These data indicate that H3N2 vaccine antigens produced during the 2016–2017 influenza season in eggs and MDCK cells (that possess K160 HA) elicit antibody responses that react poorly to current circulating H3N2 virus strains, whereas H3N2 vaccine antigens produced in insect cells (that possess T160 HA) elicit antibody responses that react efficiently to current circulating H3N2 viral strains.

The Effect of Repeat Vaccination on Anti-H3 Antibody Responses During the 2016–2017 Influenza Season. Many individuals receive influenza vaccines every year; however, recent data suggest that repetitive vaccinations may be associated with reduced antibody responses (22–24) and reduced vaccine effectiveness (25–27) during some influenza seasons. To determine whether prior vaccinations impacted the development of H3 antibodies following vaccination of donors in our study, we examined neutralizing antibody data in relation to vaccine history. Donors in our study self-reported vaccination history in the previous two seasons (2012–2013 and 2013–2014). We excluded individuals who reported having received the live attenuated vaccine in previous seasons (n = 1) and individuals who did not remember their vaccination status in either season (n = 7), and then we grouped individuals into three categories: unvaccinated both seasons, vaccinated in only one season, and vaccinated in both seasons. Interestingly, vaccination history was uncorrelated with prevaccine titers to T160 HA and K160 HA, but it was the strongest independent predictor of final antibody titers and fold following vaccination with Fluzone and Flucelvax (Fig. 4A). Conversely, antibody increases against the K160 HA were similar following vaccination with the three different vaccine antigens (Fig. 4B). These differential effects of the vaccines remained statistically significant after adjusting for age, prevaccination titer, and vaccination history (Table S3). These data indicate that H3N2 vaccine antigens produced during the 2016–2017 influenza season in eggs and MDCK cells (that possess K160 HA) elicit antibody responses that react poorly to current circulating H3N2 virus strains, whereas H3N2 vaccine antigens produced in insect cells (that possess T160 HA) elicit antibody responses that react efficiently to current circulating H3N2 viral strains.

Vaccine effectiveness was lower in younger adults compared with older adults during the 2016–2017 season (20). Interestingly, we found that younger adults had higher prevaccination titers to the egg-adapted K160 HA compared with older individuals (Spearman’s rho = 0.3, P < 0.01; raw data from three independent experiments are shown in Table S2). Following vaccination, K160 HA titers were also higher in younger adults compared with older individuals (Spearman’s rho = 0.4), even after adjusting for higher prevaccination titers to K160 HA (Table S3). We speculate that the observed age-related differences in antibody titers to K160 HA might be due to birth year-related differences in H3N2 exposure history. Antibody titers to T160 HA were lower than antibody titers to K160 HA both before and after vaccination (P < 10−16 for each, Wilcoxon rank sum test), but there were no birth year-related effects on antibody titers to T160 HA (P = 0.23 prevaccination, P = 0.33 postvaccination; Table S3).

Some individuals in our study mounted strong antibody responses against T160 HA following vaccination (Table S2). Importantly, the majority of these individuals received the Flublok vaccine that possesses T160 HA (Fig. 4A). Antibody titers against T160 HA increased approximately fourfold following vaccination with Flublok, whereas there were only minimal increases in T160 HA antibody reactivity in most individuals following vaccination with Fluzone and Flucelvax (Fig. 4A). Conversely, antibody increases against the K160 HA were similar following vaccination with the three different vaccine antigens (Fig. 4B). These differential effects of the vaccines remained statistically significant after adjusting for age, prevaccination titer, and vaccination history (Table S3). These data indicate that H3N2 vaccine antigens produced during the 2016–2017 influenza season in eggs and MDCK cells (that possess K160 HA) elicit antibody responses that react poorly to current circulating H3N2 virus strains, whereas H3N2 vaccine antigens produced in insect cells (that possess T160 HA) elicit antibody responses that react efficiently to current circulating H3N2 viral strains.

Vaccine antigens are prepared in fertilized chicken eggs, a small fraction of vaccine antigens are produced in insect cells or canine kidney cells. To account for this, we measured human antibody responses elicited by vaccine antigens prepared in eggs (Fluzone; n = 22 donors), MDCK cells (Flucelvax; n = 26 donors), and insect cells (Flublok; n = 22 donors). Importantly, the H3 antigens in the Flu zone and Flucelvax vaccines possess the egg-adapted K160 HA, while the recombinant H3 antigen in the Flublok vaccine possesses T160 HA.

In contemporary H3N2 viruses with T160 HA, even after adjusting for higher antibody titers to K160 HA both before and after vaccination (Table S3). These data indicate that H3N2 vaccine antigens produced during the 2016–2017 influenza season in eggs and MDCK cells (that possess K160 HA) elicit antibody responses that react poorly to current circulating H3N2 virus strains, whereas H3N2 vaccine antigens produced in insect cells (that possess T160 HA) elicit antibody responses that react efficiently to current circulating H3N2 viral strains.
Ferrets elicit different types of antibody responses when exposed to test. Ferrets possess different antibody responses when infected with viruses possessing (A) K160 HA or (B) T160 HA. Neutralization titers were expressed as inverse dilution of sera that reduced foci by 90%. We completed three independent experiments with each sera. Shown are geometric means from the three independent experiments. Statistical significance was determined using a paired Student’s t-test.

**Discussion**

It is unclear why there was only moderate vaccine effectiveness during the 2016–2017 influenza season, given that the 2016–2017 vaccine strains appeared to be well matched to most circulating viral strains (20). Our data suggest that a mismatch in antigenic site B of H3N2 viruses, caused by the propagation of the vaccine strain in eggs, likely contributed to this low vaccine effectiveness. When influenza viruses are passaged in new hosts, antigenic properties of hemagglutinin can change since many residues of this protein are involved with both receptor binding and antibody recognition (28–31). This is not the first example of antigenic properties of influenza viruses being altered by egg adaptations. In 1978, Kilbourne (32) showed that there were two antigenically distinct viruses in the 1976 X-53 swine influenza vaccine and that these viruses had different growth properties in chicken embryos and canine kidney cells. In 1983, Webster and coworkers (33) demonstrated that influenza B viruses acquire mutations that alter antigenicity when propagated in chicken eggs. Egg adaptations resulting in antigenic changes have also been reported for H3N2 viruses (34–37). It has been proposed that H3N2 egg adaptations contributed to low vaccine effectiveness during the 2012–2013 influenza season (38), although this remains controversial (39).

Nonetheless, the majority of influenza vaccine antigens continue to be prepared in eggs (4). One solution to the problem of egg adaptations is to simply produce influenza antigens via a baculovirus system or in cell culture. The baculovirus system seems particularly well suited to avoid adaptive mutations; however, there are also potential problems with this approach. HA antigens prepared in insect cells are glycosylated with less complex sugars compared with mammalian cells (40), and it is possible that this affects antigenicity. In our study, some individuals vaccinated with baculovirus-prepared Flublok (that possessed T160 HA) mounted strong antibody responses that effectively neutralized a virus possessing an HA with a glycosylated antigenic site B; however, some Flublok-vaccinated individuals mounted antibody responses that reacted poorly to viruses with T160 HA.

It is important to note that there is more HA antigen in Flublok vaccine formulations compared with conventional egg-based vaccine formulations. It is possible that the increased amount of antigen in Flublok contributed to higher antibody responses against viruses that possess T160 HA; however, this is likely not the case since compared with the other vaccines tested, Flublok did not induce a significantly greater response to K160 HA. Therefore, Flublok did not generate an overall higher antibody response in our study, but rather an antibody response that was better able to recognize viruses possessing T160 HA. This is interesting in light of a recent study demonstrating that the Flublok vaccine elicited more protective antibody responses in older adults compared with egg-based vaccines during the 2014–2015 season (41). This warrants further investigation because there was a large H3 antigenic mismatch during this season (17). It is possible that the Flublok vaccine elicited antibodies of different specificities compared with egg-based vaccines during the 2014–2015 influenza season, perhaps to epitopes involving conserved residues of the HA receptor binding pocket.

Cell culture-expressed HA antigen might also avoid problems associated with egg adaptations. The 2017–2018 recommended H3N2 component is the same as the 2016–2017 recommended H3N2 component, but, starting this year, vaccine manufacturers that prepare antigens via cell culture systems will be allowed to use viral strains that have not been previously adapted to grow in eggs (https://www.cdc.gov/flu/protect/vaccine/cell-based.htm). H3N2 vaccine strains grown in this new system may possess HA with a glycosylated antigenic site B, although mutations that abrogate this glycosylation site have been previously reported upon serial passage of clade 3C.2a H3N2 viruses in MDCK cells (21).

Interim reports indicate that the 2016–2017 vaccine protected younger individuals from H3N2 infection less effectively compared with older individuals (20). In our study, we found that younger adults were more likely to produce antibodies that efficiently neutralized K160 HA but weakly neutralized T160 HA after vaccination. These age-related differences may be due to...
differences in prior exposure histories. It is clear that prior influenza exposures can affect how an individual responds to new antigenically distinct viral strains (42). For example, individuals exposed to the 2009 pandemic H1N1 strain produce antibodies that target epitopes that are conserved in previously circulating seasonal H1N1 viral strains (43–46), particularly viral strains that they were likely exposed to in childhood. Early childhood infections also likely impact susceptibility to pandemic influenza virus strains. Gostic et al. (47) recently demonstrated that early childhood infections with either H1N1 or H3N2 influenza viruses are associated with protection from H5N1 and H7N9 viruses later in life, presumably through induction of cross-reactive antibodies against epitopes that are conserved between these different viruses. Further studies should explore if early childhood antigenic “ imprinting” with different H3N2 viral strains affects the specificity of antibodies elicited by seasonal influenza vaccination.

Vaccine effectiveness against H3N2 is often low (2), especially among repeat vaccinees (25–27). Antigenic mismatch due to incorrect strain selection is an established cause of low vaccine effectiveness (2). Future studies should address whether egg-adapted mutations constitute another form of antigenic mismatch that alters vaccine effectiveness in other influenza virus seasons. Antigens with egg-adapted mismatches might recall pre-existing immune responses, including responses to previous vaccines. A major effort should be made to develop and utilize new systems that produce influenza antigens that are not dependent on egg or cell culture-adaptive mutations. Antigens that do not possess adaptive mutations will likely offer better protection against influenza virus strains that circulate in the human population.

**Materials and Methods**

**Viruses.** The HA gene of a representative clade 3C.2a H3N2 virus (A/Colorado/15/2014) was cloned into the vector pHW2000, and the T160K mutation was introduced to remove the predicted glycosylation motif. Viruses containing H3 and N2 genes with A/Puerto Rico/34/1934 internal genes were rescued into 293T and MDCK-SIAT1 cells. Transfection supernatants were collected 3 d after transfection and stored at −80 °C for use in neutralization assays. HA glycoproteins were transferred to a nitrocellulose membrane (Thermo Fisher Scientific). Blots were probed using anti-HA tag primary antibody clone HA-7 (product number 59658; Sigma-Aldrich) and an anti-mouse secondary antibody (product number 926-32212; Licor). To structurally model the N158 glycosylation introduced by T160, a K160T mutation was introduced into the cryostructure (PDB ID code 4OS5) using the program PyMol. Basic N-linked glycans were added to predicted N-linked glycosylation sites in the cryostructure using the GlyProt web server: [www.glycosciences.de/modeling/glyprot/](http://www.glycosciences.de/modeling/glyprot/).

**Glycosylation Western Blotting and Glycan Modeling.** Viruses possessing T160 and K160 HA were expanded on MDCK-SIAT1 cells and then concentrated at 20,000 rpm (72,128 × g) for 1 h at 4 °C using an SW-28 rotor (Beckman Coulter). The amount of HA in each sample was normalized by ELISA with a human anti-HA monoclonal antibody for subsequent SDS/PAGE and Western blotting. To cleave N-linked glycans, samples were treated with PNGase-F (New England Biolabs) under reducing conditions. PNGase-treated and untreated samples were run on SDS/PAGE gels (Thermo Fisher Scientific) and transferred to a nitrocellulose membrane (Thermo Fisher Scientific). Blots were

**Human Monoclonal Antibody Binding.** Human monoclonal antibodies were previously isolated from donor peripheral blood mononuclear cells following vaccination with the 2010–2011 influenza vaccine (48). ELISA plates were coated overnight at 4 °C with a 2009 virus (A/Victoria/210/2009; the H3N2 component of the 2010–2011 influenza vaccine), a glycosylated 2014 clade 3C2.a virus with T160 HA, or an unglycosylated 2014 clade 3C2.a virus with K160 HA. ELISA plates were blocked with a 3% (wt/vol) BSA solution in PBS for 2 h. Plates were washed three times with PBS containing 0.1% Tween 20, and serial dilutions of each monoclonal antibody in ELISA buffer (1% BSA (w/vol) in PBS) were added to plates in 1% BSA in PBS. After 2 h of incubation, plates were again washed and a peroxidase-conjugated anti-human secondary antibody (product number 109-036-098; Jackson ImmunoResearch) diluted in ELISA buffer was added. A 1-h incubation, plates were washed, and a 3,3′,5,5′-tetramethylbenzidine (TMB) substrate (product number 50-00-03; Seracare) was added. The TMB reaction was stopped by adding HCl, and absorbance was measured using a plate reader. One-site specific binding curves were fitted in Prism software and the maximal binding against the 2009 vaccine strain was calculated for each monoclonal antibody. After background subtraction (from ELISA plates that did not have coated virus), binding of each monoclonal antibody relative to the vaccine strain was determined using the absorbance values at the lowest concentration of antibody that gave more than 90% of maximal binding against the vaccine strain. Equivalent coating of each virus was checked using the stalk-reactive DY-2F04 monoclonal antibody.

**Ferret Sera.** Ferret antisera were prepared at the Association for Assessment and Accreditation of Laboratory Animal Care-accredited company Noble Life Sciences using protocols approved by the Noble Life Sciences Institutional Animal Care and Use Committee. Naïve fitch ferrets were intranasally infected with 2 × 105 PFU of virus expressing either K160 HA or T160 HA. Serum samples were collected before infection and 28 d following infection. Sera used in antigenic assays were treated with receptor-destroying enzyme (Denka Seiken) for 2 h at 37 °C, and the enzyme was then heat-inactivated at 55 °C for 30 min.

**Human Sera.** Experiments using human sera were conducted with the approval of the University of Rochester and University of Pennsylvania Institutional Review Boards. Informed consent was obtained for all individuals enrolled. Serum samples were collected at the University of Rochester before and 28 d following vaccination. Serological experiments were completed at the University of Pennsylvania using deidentified samples. Before assays, sera samples were treated with receptor-destroying enzyme (Denka Seiken) for 2 h at 37 °C, and the enzyme was then heat-inactivated at 55 °C for 30 min.

### Table S3

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### Table S4

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FRNT Assays. Serum samples were serially diluted in 96-well round-bottom plates containing serum-free media. Approximately 200 focus-forming units of reverse-gene engineered infectious variant of each virus were added to each dilution step and the virus-sera mixtures were incubated at room temperature for 1 h. The virus-sera mixtures were then added to confluent monolayers of MDCK-SIA1T1 cells and incubated at 37 °C for 1 h. After incubation, cells were washed with serum-free media and an overlay medium of serum-free media containing 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (Hepes) (Thermo Fisher Scientific), gentamycin (Thermo Fisher Scientific), and 0.5% (wt/vol) methylcellulose (Sigma-Aldrich) was added to cells. Infected monolayers were incubated for 18 h, after which the overlay medium was aspirated and the cells were fixed and permeabilized with ice-cold methanol–acetone [1:1 (vol/vol)]. Infected monolayers were stained with anti-NP monoclonal antibody IC5-187 (product number NR-43899; BEI Reagent Resources) and an anti-mouse peroxidase-conjugated secondary antibody (product number 855563; MP Biomedicals). Plates were washed with serum-free media and an overlay medium of serum-free media containing 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (Hepes) (Thermo Fisher Scientific). Infected monolayers were washed with serum-free media and an overlay medium of serum-free media and methylcellulose (Sigma-Aldrich) was added to cells. Infected monolayers were incubated for 1 h. Following staining, plates were imaged and foci were quantified using an ELISPOT reader (Cellular Technologies Limited). FRNT titer were reported as the reciprocal of the highest dilution of sera that reduced the number of foci by at least 90%, relative to control wells that had no serum or monoclonal antibody added. Undetectable titters were assigned a value of 10. All FRNT assays were repeated three times on separate days.

Models of Initial and Final Antibody Titers and Fold Responses. Three replicate measurements of pre-vaccination and post-vaccination antibody titers from each serum sample (Table 5) were geometrically averaged, and the geometric mean titers were used for model fitting. Linear models that included age and vaccination history (receipt of trivalent inactivated vaccine in neither, only one, or both of the last 2 y) were fitted to log antibody titers as continuous and factor variables, respectively. Models were similarly fitted to log, post-vaccination antibody titers, also including log, pre-vaccination titters and vaccine group (Flublok, Flucelvax, and Fluzone) as continuous and factor variables, respectively. Likelihood maximization was performed with the lm package in R, version 3.3.1. Code for the analysis is available at https://cobeylab.github.io/H3N2_glycosylation/.  

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