Hemagglutinin Glycan Clock Guides Human Influenza A Virus Evolution

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ABSTRACT

Glycosylation of the influenza A virus hemagglutinin (HA) globular domain greatly influences viral antigenicity, receptor binding, and innate immunity. Gel electrophoresis and bioinformatics analysis reveals that circulating human H1 and H3 HA glycans largely evolve in a clock-like progression according to three simple rules. This glycan clock predicted the 2015-2016 selective sweep of pH1N1 by strains with an added glycan at residue 179. A limit on glycan shielding of the HA head dictates the ultimate fate of circulating strains, enabling our prediction that circulating H3 HA strains will either undergo a glycan readjustment after 9 years, or be supplanted by a novel pandemic virus.

INTRODUCTION

easonal influenza A virus (IAV) annually sickens millions and kills hundreds of thousands of people (WHO, 2003). IAV remains endemic despite high levels of adaptive immunity in nearly all humans after early childhood. IAV persists due to the rapid selection of mutants with amino acid substitutions in the viral hemagglutinin (HA) glycoprotein that enable escape from neutralizing antibodies (Abs). HA is a homotrimeric glycoprotein with a globular head domain resting atop a stalk that anchors HA to the virion surface. The head has a receptor binding site that attaches virions to cell surface terminal sialic acid residues, initiating infection. Ab binding to the head neutralizes virus by blocking attachment or the conformational alterations required for HA-mediated membrane fusion. Understanding Ab-driven HA evolution is essential to improving influenza vaccination, which at best, offers limited protection from infection (CDC, 2017; Couch and Kasel, 1983).

As newly synthesized HA enters the endoplasmic reticulum of IAV infected cells, oligosaccharides are attached to Asn residues present in the motif Asn-X-Ser/Thr-Y,

where X/Y are all amino acids except Pro. Attached glycans can be high-mannose glycans or larger, more complex branched glycans (Keil et al., 1985; Khatri et al., 2016). HAs possess multiple highly conserved glycans on the stem domain crucial for HA folding and oligomerization (Daniels et al., 2003). Glycosylation of the head is much more variable, as evolving HAs add, retain, and occasionally lose potential head glycosylation sites (An et al., 2015; Sun et al., 2011; Vijaykrishna et al., 2015). Head glycans can promote viral fitness by shielding virus from antibody binding and tuning receptor affinity and specificity (Job et al., 2013; Medina et al., 2013; Sun et al., 2013b; Vigerust et al., 2007). Conversely, glycans can deleteriously modulate receptor binding, enhance viral neutralization by innate immune lectins, or increase stress in the endoplasmic reticulum during translation (Alymova et al., 2016; Das et al., 2011; de Graaf and Fouchier, 2014; Hrincius et al., 2015; Park et al., 2016; Yang et al., 2015). Glycosylation changes are generally not correlated with changes between antigenic clusters, in part, because glycan addition typically lowers HA receptor avidity (Aytay and Schulze, 1991), complicating binary assays such as hemagglutination inhibition (Das et al., 2011). Consequently, they are typically unaccounted for in molecular evolution studies using HI-based analytic tools such as antigenic cartography (Blackburne et al., 2008; Sun et al., 2013a).

HA gel electrophoresis suggests rules for glycan evolution

Studies of HA glycosylation evolution rely nearly exclusively on bioinformatic prediction of glycan addition. NetNGlyc, the state of the art algorithm (Gupta, 2004), imperfectly predicts N-linked glycosylation (Khatri et al., 2016). We therefore used gel electrophoresis to monitor glycan addition to HAs from 72 egg-grown H1N1 and H3N2 strains based on decreased HA migration (**Fig. 1**, **fig. S1-S4**). HA migration strongly correlates with the number of computationally predicted head glycosylation sites (Pearson coefficient r = 0.96, P < 0.01) (**Fig. 1A**). H1 adds an apparent 3.9 ± 0.1 kDa [$R^2 = 0.89$] per glycan, suggesting the majority of such glycans are highly branched. The average H3 glycan, by contrast, is smaller (3.0 ± 0.2 kDa) ([$R^2 = 0.90$]), suggesting heterogeneity in glycan mass or incomplete glycosylation. This size difference between H1 and H3 HAs is consistent with mass spectrometry characterization showing that H1 glycans skew to larger more complex structures, while H3 glycans at several sites are majority highmannose (Khatri et al., 2016; Sun et al., 2013b).

Importantly, for H1 and H3 HAs, the total glycan mass added to the head during evolution in humans is, respectively, 15.5 kDa and 15 kDa. This is consistent with the idea of an upper limit of glycan shielding for H1 and H3 HA head domains totaling to ~20 kDa (5 glycans for H1, 7 for H3). This is not strictly due to the replicative capacity of IAVs: H1 HAs can accommodate at least 8 head glycans while maintaining *in vitro* fitness (Eggink et al., 2014). Rather it points to glycan-based fitness costs *in vivo*, possibly due innate immune mechanisms (Tate et al., 2014).

Plotting HA migration versus time reveals the history of seasonal H1N1 (sH1N1) glycosylation (**Fig. 1B**, triangles). First-wave sH1N1 (1933-1957) HAs behaved differently than the second-wave sH1N1 representatives (1977-2008). During the first-wave, HA increased by an average of 3.9 kDa (or one glycan) every 5 years (slope = 0.69 kDa/year, $R^2 = 0.72$). The H1N1 strains that re-emerged after the 1977 introduction of H1N1 exhibited similar HA migration to the 5-head glycan A/Fort Leonard Wood/1951 (inside blue box, **Fig. 1B**). Rather than increasing in number, electrophoresis confirms the NetNGlyc prediction that head glycans were maintained until decreasing by one unit after 1991.

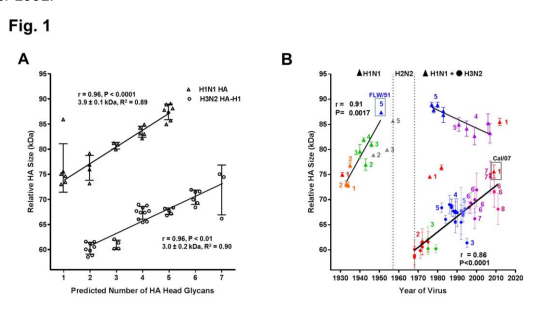


Fig. 1. Quantitative survey of glycan evolution in IAV. (A) Relative HA size, measured by migration rate through a gel, correlates with predicted number of head glycans. H1N1s (triangles) add 3.9 ± 0.1 kD per glycan, and 15.5 kD total 1 to 5 glycans. H3N2s (HA1, circles) add 3.0 ± 0.2 kD per glycan and 15 kD total in increasing from 2 to 7 glycans. **(B)** HA size over time for H1N1s (triangles) and H3N2s (HA1, circles). Strains are color-coded to match glycan groups in **fig. S2.** Numbers of head glycans per strain are noted next to individual points. A glycan is added on average every 5 years for sH1N1 (3.9 kD) and every 8 years for H3N2 (3.0 kD). The strain most related to sH1N1 reintroduced in 1977, A/Fort Leonard Wood/1951, is boxed and labelled in blue. The pH1N1 strain, A/California/07/2009, is boxed and labelled in black. Pearson coefficient of correlation (r) and its P value are shown on both graphs. Error bars are standard error of the means from n = 3-5 blots.

Bioinformatic analysis supports rules of glycan evolution

The second-wave sH1N1 began in 1977 with the re-introduction of a strain genetically similar to A/Fort Leonard Wood/1951, possibly due to a flawed vaccine trial (Rozo and Gronvall, 2015) (**Fig. 2**). With 5 head glycans, reemerged sH1N1 reached our proposed H1 head-glycan ceiling compatible with competitive human circulation. Indeed, sequences predicted to have 6 glycans are unusual over the century of H1 evolution (1918-2017), except in 1986 and 1987, when they competed among H1 strains (**fig. S5A**). Rather than adding glycans, reemerged sH1N1 viruses exhibit glycan adjustments at roughly 10 year intervals. In 1987, N71 pH1N1 (numbering convention is shown in Table S1) replaces N172, as N144 toggles to N142, which though adjacent to 144, is oriented to shield different epitopes. In 1997, N286 is lost, and this glycan configuration remained constant until sH1N1 became extinct in 2009.

Turning to H3 glycan evolution (**Fig. 4**), original (1968) pandemic H3 strains have two head glycans. Six years after the pandemic, H3N2 added a third glycan, N126. Six years later, it added another glycan at either N246 or N122. The virus then persisted 12 years with 4 glycans, suggesting a functional limitation to adding a fifth glycan. In 1993, an antigenically distinct H3N2 swept the globe; this virus added a glycan at N276, which is adjacent to a disulfide-bonded cysteine. This 5-glycan H3N2 disappears in 3 years, and by 1998 is completely replaced by HAs with 2 added glycans, N122 and N133, for a net increase of one glycan. It takes another 6 years for complete replacement by strains that have a seventh glycan, N144.

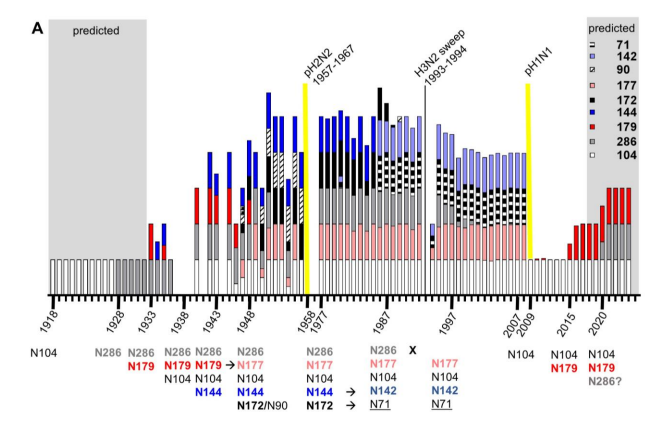
With 7 glycans, H3 HAs had increased 15 kDa from the original 1968 strain. It is apparent from the global human influenzome (GHI) that H3 HAs with more than 7 glycans are unfit for human circulation—8-glycan strains make up <1% of H3N2 sequences in any year (**fig. S5B**). Unable to add a glycan to escape immune pressure, 12 years after reaching the glycan limit, H3N2 underwent a selective sweep that toggled the glycan from N144 to N158 by 2015 (**fig. S6A**).

Influenza glycan clock algorithm

Based on these observations we propose 3 rules that explain, albeit imperfectly, the evolution of HA glycans:

- 1. Glycans are limited on the HA head to 5 and 7 glycans, respectively, for H1 and H3 HAs, with a total mass of ~20 kDa due to preferred use of complex (H1) vs. simple (H3) glycans.
- 2. If HA can add a glycan and retain high transmissibility, it will add one after 5 years for H1 or 6 years for H3.
- 3. If a glycan addition cannot occur, after 10 years for H1, or 12 years for H3, either a glycan switch will occur or a new pandemic strain will replace it.

Fig. 2



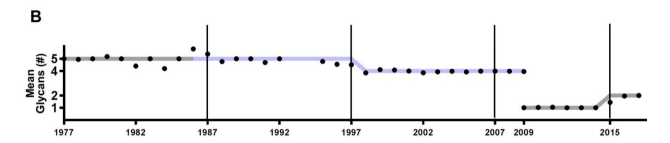


Fig. 2. H1 Glycan Clock 1918-2022. Stacked bars correspond to the percentage of all human H1N1 sequences from a given year containing a glycan listed in the legend. Dates of glycan transition are written on the timeline below along with the glycosylated residues. Projected values for strains from 1918-1933 and 2017-2022 are shown in shaded regions. Years with no data are left blank. Sequence counts for each year are shown in fig. S6. Pandemics are denoted by yellow lines. (B) The mean predicted number of glycans among all sequences from each year are shown as black circles. This data is laid atop the number of glycans predicted from the glycan clock algorithm (grey line, light blue line indicates glycan transition from sequences containing N144/N172 to N142/N71). Black vertical lines denote predicted years for glycan evolution.

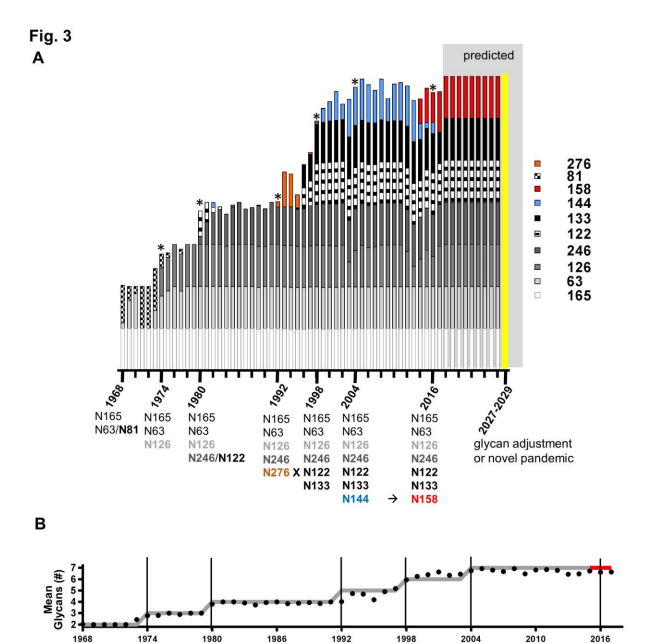


Fig. 3. H3 Glycan Clock 1968-2029. (A) Stacked bars correspond to the percentage of all human H3N2 sequences from a given year containing a glycan listed in the legend. Dates of glycan transition are written on the timeline below along with the glycosylated residues, as well as denoted by asterisks. Projected values for strains from 2017-2029 are shown in shaded regions. A potential pandemic is denoted by yellow line. **(B)** The mean predicted number of glycans among all sequences from each year are shown as black circles, which are displayed atop the number of glycans predicted from the glycan clock algorithm (grey line, red line indicates glycan transition from sequences containing N144 to N158). Black vertical lines denote years of glycan evolution.

Predicting future influenza evolution events

Using these rules, we can predict future IAV glycosylation evolution (**Fig. 2** and **3**, grey boxes). H3N2 toggled N144 to N158 over the 2014-2015 season. According to Rule 1, H3N2 is limited to 7 glycans. By Rule 3, during the 2027-2029 seasons, H3N2 should either exhibit another glycan adjustment or be replaced by a new pandemic strain.

In 2009, pandemic H1N1 (pH1N1) rapidly replaced sH1N1 in the GHI. A single head residue is predicted to be glycosylated (N104). Indeed, the HA of a representative early strain, A/California/07/2009 (**Fig. 1B**, inside black box), co-migrated with strains from the 1930s and two other swine-origin human strains, A/New Jersey/8/1976 and A/Memphis/4/1982, each with a single predicted head glycan site (**Fig. 1B**, red triangles).

Based on the addition of a globular-domain glycan to first wave H1 strains every 5 years from 1933 to 1951, we predicted that pH1N1 viruses should add a glycan by 2015. Indeed, identifying all potential HA head glycosylation sites in pH1 sequences in the Flu Database (FluDB) revealed a second predicted glycan based on a S to N change at residue 179 (**Fig. 4A**). N179 viruses are present at low frequencies in each season since pH1N1 introduction (**Fig. 4B**), but over 15 months during 2015-2016, N179 replaced all previously circulating strains in the pH1N1 GHI (**fig. S6B**).

How did the N179 strains come to selectively sweep the human pH1N1 GHI? We identified the earliest genetic precursor "sweeper strains" (SWSs) of the N179 strains in the FluDB by aligning pre-2016 N179-containing strains with Clustal Ω software to generate an evolutionary tree. The first pre-SWSs appeared almost simultaneously in January 2015 in Iran (4 strains), Turkey (1 strain), and Taiwan (1 strains). In addition to an additional glycan site at N179, all SWSs carried the I233T substitution. This combination of I233T and N179 is not found in prior strains.

The A/California/07/2009 HA structure provides a ready explanation for the co-selection. Residues 233 and 179 are in close proximity across the trimer interface (**Fig. 4C**), with the change from bulkier and more hydrophobic lle to Thr likely to facilitate the addition of the large and polar glycan on the other side of the trimer interface. Notably, in H3 strains with a N179 glycan, the residue analogous to 233 in pH1N1 is also a small polar residue (Ser).

As the first 3 glycans added to sH1N1 were N104, N179, and N286 (see strain A/Hickox/JY2/1940), we predict that N286 will be present in the next glycan selective sweep strain, which based on Rule 2, should occur during the 2020-2021 season.

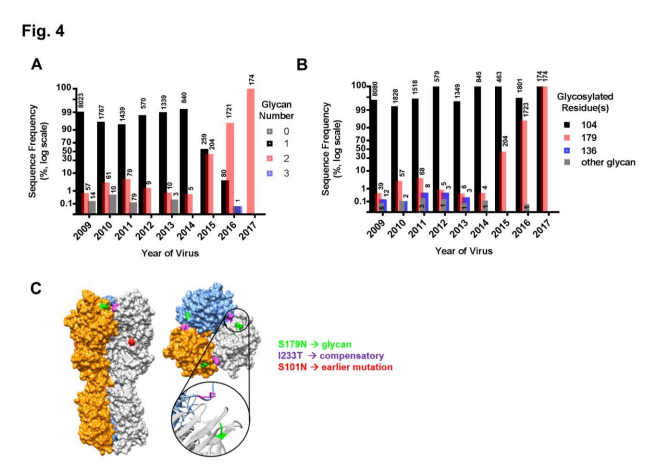


Figure 2. Glycan clock predicts recent pH1N1 glycan addition. (A) Frequency of human pH1N1 sequences in FluDB by number of predicted glycans in each calendar year. The number of sequences represented by each bar is shown on top of bar. Until January 2015 >90% of sequences had 1 glycan (black). Starting in 2015, strains with 2 glycans swept (red). (B) Frequency of the individual residues glycosylated in human pH1N1 sequences. N104 predominates throughout (>95%, black). N179 (red) and N136 (blue) appear at low frequencies until 2015 when strains having both N179 and N104 extinguish N104-only strains. N179 strains are found in every year. Rarely, in addition to N104 a residue other than N179 or N136 has a potential glycosylation site (grey). (D) A/California/07/2009 (PDB: 3LZG) hemagglutinin crystal structure with N179 highlighted in green and the compensatory mutation I233T in purple adjacent across the trimer. Inset shows magnified view of the two mutated residues.

CONCLUSION

It has been known for some time that IAV maintains circulation in humans, in part, by altering the number and type of N-linked glycans on the globular head of HA. After measuring the relative size of HA within a historical panel of IAVs we hypothesized: (1) there is a functional limit to the number of HA-head glycans IAV can add while still retaining fitness in the human population; and (2) glycan evolution in IAV occurs at a

regular rate. When we tested these hypotheses, bioinformatically against the set of all human influenza sequences, clear patterns emerged, leading to development of the Influenza Glycan Clock algorithm. This algorithm predicted the 2015-2016 selective sweep of pH1N1 by strains with a new glycan at N179.

Going forward, we envision this algorithm will permit us to predict the timeline and trajectory of IAV evolution from their pandemic introduction to eventual replacement. We anticipate this discovery will have important implications for improving pandemic preparation and vaccine development.

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SUPPLEMENTAL MATERIAL

Material and methods fig. S1-S6 Table S1

MATERIALS AND METHODS

Virus preparation and purification

We propagated each virus by infecting 5, 10-day old embryonated chicken eggs with 100µL of viral stock diluted 1:2000 in PBS. Infected eggs were incubated at 34.5°C and 65% humidity for ~41h. We first clarified the allantoic fluid (AF) by centrifuging 4500 x g for 20min, then pelleted the virus by centrifuging for 2h at 26,000 x g. After incubating the viral pellet in 1.7mL PBS overnight at 4°C, we layered the virus onto a discontinuous 15%-60% sucrose gradient and spun at 34,000 x g for 2h. We collected virus at the gradient interface and spun it down a final time in PBS at 34,000 x g for 2h. Virus was resuspended in 100µL PBS and stored as milky opalescent suspensions long term at 4°C. Hemagglutination titers for each virus were measured from neat AF. A complete list of viruses and their origin is found in **File S1**. Total protein content of each virus prep was determined by Coomassie blue-based assay performed per manufacturer instructions [Bio-Rad, DC Assay].

Allantoic fluid (AF) from eggs infected with 93 out of the 97 viruses we attempted to grow showed hemagglutination activity (HAU), indicating viral presence. We collected pure virus from all viruses in amounts ranging from 20-6920 μ g. Sham infection and purification of AF from control eggs did not produce enough protein for detection, and ran cleanly on a gel (10 μ L of a 100 μ L prep, Lane #29, **fig. S1**).

To determine the identity of individual bands on the Coomassie gel, we grew H1N1/H3N2 reassortant viruses A/HK/68 (HK), A/PR8/MCa (PR8), X31 (HK HA and NA, PR8 background), J1 (HK HA, PR8 background), and P50 (PR8 HA, HK background). These viruses were run together on the same gel and immunoblotted with appropriate antibodies to identify which bands corresponded to which proteins on via Coomassie blue staining (**fig. S2**).

Protein gels and immunoblotting

We mixed purified virus (1 µg protein) with 4× NuPage loading buffer (Invitrogen), and boiled for 10 min at 96°C. We electrophoresed samples with Chameleon Duo Li-Cor ladder on 4–12% Bis-Tris Gels (Invitrogen) at 200 V for 55 min. To visualize proteins, we fixed gels for 10 min with 10 ml 10% acetic acid and 50% methanol, shaking at RT. After removing fixative, we added 10 ml GelCode stain (Pierce) and shook for 30 min at RT, then destained the gels with water overnight. For immunoblotting, we transferred proteins from gels to PVDF membranes with the iBLOT at P3 setting for 7 min. We blocked membranes for 30 min at room temperature (RT) StartingBlock (Thermo). After incubating with primary Ig or sera for 1 hr at RT, and washing 5x for 5 min each in TBST (10 mM Tris, 150 mM NaCl, 0.1% Tween-20), we added secondary Ig, repeating the washing step incubation. We imaged blots and Coomassie gels on a Li-Cor Odyssey. Band signals and molecular weights were calculated using Li-Cor's ImageStudio software.

Glycoprotein molecular weights were normalized for gel distortion, commonly called "smiling," by adjusting each glycoprotein molecular weight by the average amount that the M1 and NP molecular weight for each virus varied from the mean molecular weight of M1 and NP proteins on each gel (fig. S3). Linear regression, Pearson coefficient, and P-values were calculated using PRISM software.

Influenza sequences and N-glycosylation prediction

Human HA protein sequences were retrieved from the NIAID Influenza Research Database (IRD) (Zhang et al., 2017) through the web site at http://www.fludb.org (accessed April 20, 2017). Sequences with identical strain names were retained only if the majority of members had identical sequences. Sequences were aligned to A/California/04/2009(H1N1) (FJ966082) with MAFFT v7.305b, and removed if indels existed in relation to the reference or if any ambiguities existed within the HA head. The amino acid numbering of sites important for this work with both H1 and H3 numbering conventions is shown in **Table S1**. Potential glycosylation sites were identified by searching for the NX[S/T]Y motif where X/Y were any amino acid other than proline or by using NetNGlyc 1.0 artificial neural network-based prediction software (Gupta, 2004). For the quantitative analysis in **Fig. 1** and **fig. S4**, residues with positive NetNGlyc scores were considered glycosylated "N+", and residues with only one negative mark were recorded as "N-". Both conditions were treated as having a glycan present. Glycan predictions with "N--" were counted as negative.

pH1N1 Sweeper Strains bioinformatic analysis

All pH1N1 in FluDB from 2009 to present day were collected and annotated with collection country, date, and glycosylation status. Sequences glycosylated at N179 from 2009-2015 were aligned by ClustalΩ (Sievers et al., 2011). An unrooted phylogenetic tree of all 365, N179-glycosylated sequences was made using FigTree (Rambaut and Drummond, 2009). The earliest strains in the same clade as the SWSs were identified as: A/Shiraz/4/2015, A/Shiraz/3/2015, A/Taipei/0021/2015, A/Adana/08/2015, A/Shiraz/6/2015.

To identify any additional mutations occurring with N179 we identified all sequences with >99% sequence similarity to the 6 earliest SWSs using USEARCH (max ~17nt differences) (Edgar, 2010). This represented 15% of all FluDB sequences. An unrooted phylogenetic tree of a downsampled set of 370 sequences representing pre-SWS and SWSs (as well as outgroup A/California/07/2009) was aligned in Clustal Ω is shown to the right. From this alignment, the two conserved amino acid changes that define SWSs, S179N and I233T were clearly seen.

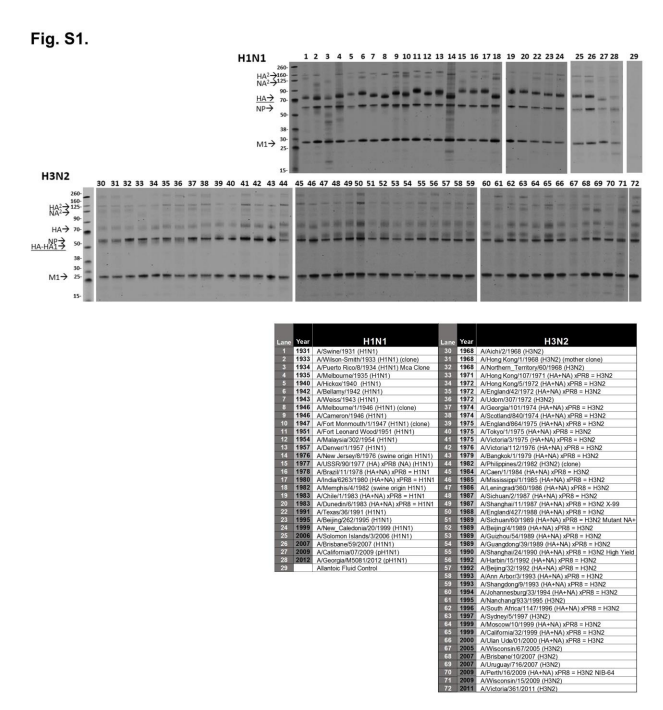
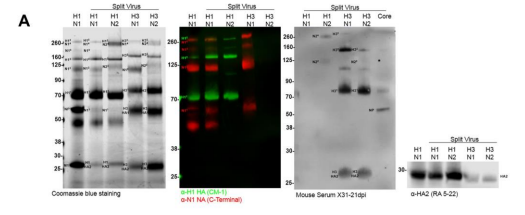
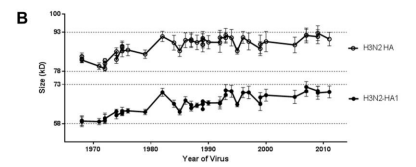


Fig. S1. SDS-PAGE of purified virus. Molecular weight standards are labelled. Representative blots are shown. Lane identifications with reassortant status are shown below blots.







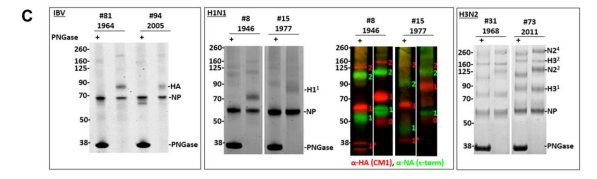


Fig. S2. Identification of viral proteins in Coomassie blue stained gels. (A) Purified whole PR8 or reassortant, detergent split viruses were stained with Coomassie blue; probed with monoclonal antibodies against HA1 (CM1), N1 (C-terminal), and HA2 (RA-5 22); or convalescent sera from a H3N2 reassortant infected mouse. **(B)** H3N2 HA monomer and H3N2-HA1 band sizes correlate almost perfectly. As the resolution was better for H3N2-HA1, these were used for analysis. **(C)** PNGase treatment of viruses shows deglycosylation of glycoproteins for representative viruses. Coomassie stained gels and monoclonal antibodies against H1 (CM1) and N1 (C-terminal) for H1N1.

Fig. S3.

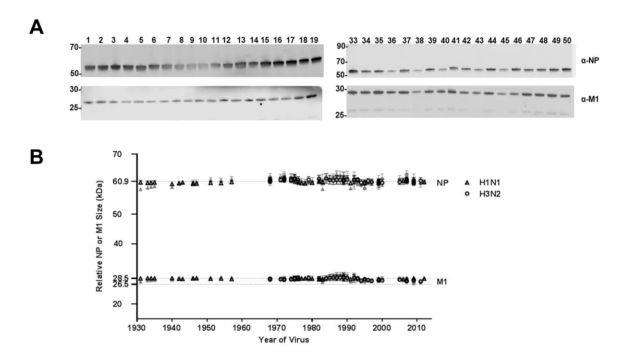
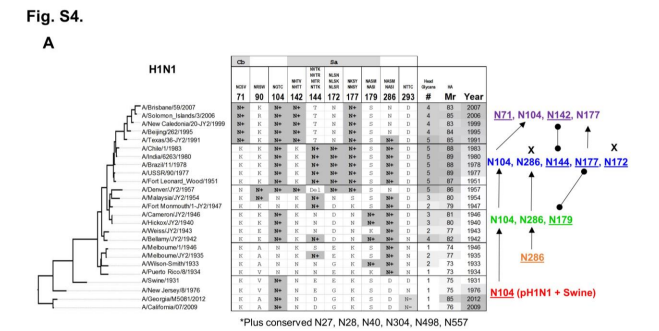
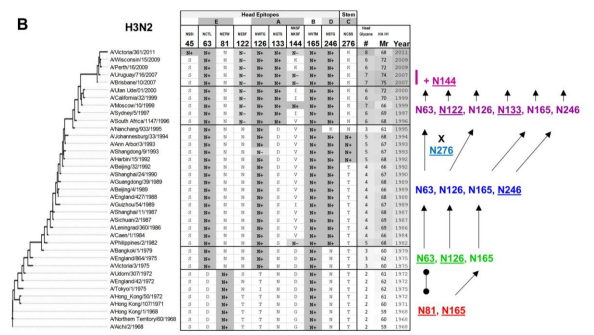


Fig. S3. NP and M1 proteins serve as internal ruler. (A) Western blots of NP and M1 for representative viruses. **(B)** Quantitation of NP and M1 relative size from SDS-PAGE. Raw data (grey) was normalized to account for gel distortion (open black symbols). Average size of each protein is shown on the Y-axis. Error bars are standard error from n = 3-6 blots.

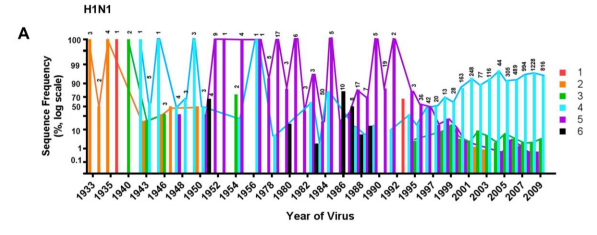




*Plus conserved N8 (except Hong Kong/50/1972 + Hong Kong/107/1971), N22, N38, N285, N483

Fig. S2. Glycan groups of IAV viral panel. (A) Phylogeny of H1N1 or **(B)** H3N2 viruses and NetNGlyc predicted glycosylation status of HA Asn (N+ or N-), the substituted residue present, or a deletion (Del). The total number of head glycans, average HA size, and year of virus collection is shown for each virus. The glycosylation progression is shown color coded to match individual viruses in **Fig. 1B**. New glycosylation sites are underlined. Arrows connect conserved glycosylation sites and barbells connect glycans in the same epitope, but not same site. Lost glycosylation sites are noted with an X. Conserved glycans among all viruses are listed below the table.





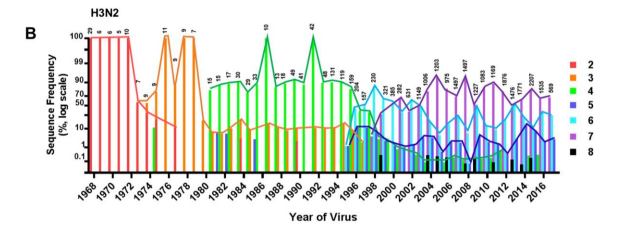


Fig. S5. Frequency of total sequences by number of glycosylation sites. (A) Percentage of H1N1 or (B) H3N2 sequences in FluDB with indicated number of predicted head glycans. Above each lane are total number of sequences in FluDB from that year. Y-Axis is log scale. Sequences above the glycan limit 6 for H1N1 and 8 for H3N2 are shown in black.

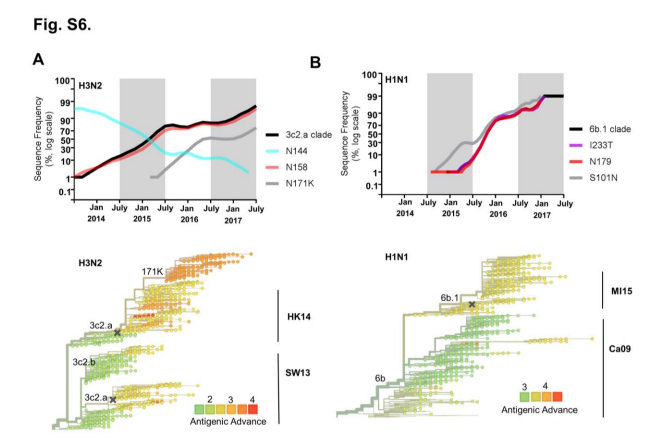


Fig. S6. Major IAV clades track with glycan evolution. Using nextflu.org we tracked the most recent glycan changes for **(A)** H1N1 and **(B)** H3N2 viruses in terms of frequency of the major clade (black) and other mutations of interest (top). On bottom, phylogenic trees are colored according to antigenic advance measured as log₂ HI titer distance from the root of the tree. Vaccine strains are marked with an X. Antigenic clusters are labelled to the right.

Table S1. Relevant amino acid residues with H1 and H3 numbering. Calculated using the "HA subtype numbering conversion tool (beta)" on FluDB. Numbering in our text starts from the initial HA methionine (M=1).

In Paper	M=1	H1	Н3
N27	27	10	20
N28	28	11	21
N40	40	23	33
N71	71	54	63
N90	90	73	82
S101N	101	84	92
N104	104	87	94
N136	136	119	125.1
N142	142	125	129
N144	144	127	131
N172	172	155	158
N177	177	160	163
N179	179	162	165
I233T	233	216	219
N286	286	269	271
N304	304	287	289
N498	498	481	483
N557	557	HA2 213	541