Abstract

Seasonal influenza virus infections can cause significant morbidity and mortality, but the threat from the emergence of a new pandemic influenza strain might have potentially even more devastating consequences. As such, there is intense interest in isolating and characterizing potent neutralizing antibodies that target the hemagglutinin (HA) viral surface glycoprotein. Here, we use cryo-electron microscopy (cryoEM) to decipher the mechanism of action of a potent HA head-directed monoclonal antibody (mAb) bound to an influenza H7 HA. The epitope of the antibody is not solvent accessible in the compact, prefusion conformation that typifies all HA structures to date. Instead, the antibody binds between HA head protomers to an epitope that must be partly or transiently exposed in the prefusion conformation. The “breathing” of the HA protomers is implied by the exposure of this epitope, which is consistent with metastability of class I fusion proteins. This structure likely therefore represents an early structural intermediate in the viral fusion process. Understanding the extent of transient exposure of conserved neutralizing epitopes also may lead to new opportunities to combat influenza that have not been appreciated previously.

Author summary

Influenza viruses cause severe respiratory infections on a global scale annually. Vaccine efforts are hampered by the virus’s naturally high mutation rate, which results in wide variation between influenza strains of the antigens that are produced and recognized by antibodies, particularly in the surface glycoprotein hemagglutinin (HA). However, broadly neutralizing antibodies (bnAbs) are a class of antibodies that develop during natural
Infected cells release mature virions, which enter the nasal mucosa and replicate to produce large numbers of new virions. A variety of respiratory and extrapulmonary symptoms are seen in human beings infected with H5N1. Laboratory studies have shown that chickens and turkeys are highly susceptible to infection by H5N1, while quails and pigeons are much less susceptible. The precise mechanism(s) by which H5N1 causes human disease is not known, but it is likely that the virus enters human beings by way of the respiratory tract. The virus is then spread to other parts of the body, possibly via the bloodstream. The symptoms of H5N1 infection in human beings vary from mild to severe, depending on the strain of the virus and the individual's immune response. In some cases, H5N1 infection can be fatal, with mortality rates ranging from 3% to 97%.
reservoirs, causing a spike in human infections in recent years [7]. Several H7-specific mAbs have been described recently, including H7.137, H7.167, and H7.169, which target highly conserved regions of the HA head adjacent to the RBS [10]. Another antibody isolated in the same study, mAb H7.5, was shown to recognize an epitope that overlaps a portion of the H7.167 epitope and neutralize similar strains of human H7 virus, including those from mallards and chickens, but which binds at a different angle [10]. The H7.5 mAb potently neutralizes human H7 strains of influenza virus, including strains isolated in outbreaks from 2003–2013 in Shanghai, the Netherlands, and New York [10].

In the course of our epitope-mapping studies of H7-specific mAbs using negative-stain electron microscopy (nsEM), we noted a striking effect of mAb H7.5 binding to HA, in which the soluble H7 ectodomain trimer falls apart. Using biochemical and structural approaches described here, we delineate the phenotypic effect of H7.5 on HA trimers. We also report several cryo-electron microscopy (cryoEM) structures of H7.5 fragment antigen binding (Fab) bound to cleaved or uncleaved H7 HA trimers. Together, these data reveal a previously unappreciated antigenic determinant on HA that, while somewhat inaccessible, may be exploited as a new target for vaccine design, given its relative conservation and structural importance. These studies also indicate that the HA trimer is likely sampling subtly different prefusion conformations that may provide clues about the early aspects of the conformational changes that accompany the fusion process.

Results

Antibody H7.5 was described in our previous study that identified several broadly neutralizing antibodies isolated from the peripheral blood B cells of donors who participated in a vaccination trial with monovalent, inactivated influenza (H7N9), A/Shanghai/02/2013 vaccine candidate (DMID13-0033) [10]. mAb H7.5 was shown to neutralize H7 viruses and exhibit strong inhibition of hemagglutination activity against a variety of H7 HAs. Preliminary analysis revealed that H7.5 recognizes multiple strains of H7 HAs with KD (dissociation constant) of less than 0.1 nM, even in the monovalent Fab form, when measured using biolayer interferometry (S1 Fig). The breadth and high affinity for the emerging H7N9 viruses made H7.5 an interesting target for in-depth structural studies. Hence, we first determined the structure of ungagged H7.5 Fab by X-ray crystallography at 2.0 Å resolution (Fig 1A, Table 1), but its complex with H7 HA could not be obtained, despite extensive screening of crystallization conditions. In parallel to the crystallization efforts, we used nsEM to characterize H7.5 Fab in complex with HA protein from different H7 strains. The difficulty in obtaining crystals of the complex became immediately apparent upon inspection of the complexes in nsEM (Fig 1B and 1C).

Reference-free 2D classification of cleaved H7 NY HA (A/New York/107/2003) in complex with H7.5 Fab showed substantial heterogeneity relative to previously observed prefusion complexes of HA bound to other head-binding antibodies [10] (Fig 1B and 1C). In nsEM 2D classes, we observed several different phenomena, including variable stoichiometries of bound Fabs and heterogeneous species that were difficult to interpret (Fig 1B and 1C). The effect of H7.5 was observed consistently with both cleaved and uncleaved forms of H7 HA from multiple strains including H7 NY HA (A/New York/107/2003), H7 Sh2 HA (A/Shanghai/2/2013), and H7 NL HA (A/Netherlands/219/2003) (S2 Fig). We therefore attempted to capture intermediate conformations of the complex by incubating the cleaved trimer with H7.5 for shorter periods of time. Indeed, a 5-minute incubation resulted in enough reliable observations to deduce that H7.5 bound to the HA1 head but induced an unfamiliar structural phenotype in our 2D classes (Fig 1B). Within a single data set, all combinations of the H7.5/H7 HA complex stoichiometry were observed (from zero to three Fabs per trimer and individual protomers
bound to H7.5 Fab) that probed the conformational landscape of HA (Fig 1B and 1C). Relative to other prefusion HA structures, the HA1 heads appeared separated, with H7.5 apparently prying them apart or stabilizing a more open form of the HA trimer (Fig 1D).

Unlike cleaved H7 HA trimers that rapidly fell apart into protomers, uncleaved H7 HA (HA0) remained in a trimeric conformation even after overnight incubation with H7.5.
Table 1. X-ray data collection and refinement statistics for H7.5 Fab.

| Data collection | H7.5 Fab |
|-----------------|----------|
| Beamline        | APS 23 ID-D |
| Wavelength (Å)  | 1.03520  |
| Space group     | P22,2,1  |
| Unit cell parameters (Å, º) | \( a = 65.5 \), \( b = 99.0 \), \( c = 208.6 \); \( \alpha = \beta = \gamma = 90 \) |
| Resolution (Å)  | 50.00–2.00 (2.03–2.00) |
| Observations    | 624,618  |
| Unique reflections | 91,327 (4,421) |
| Redundancy      | 6.8 (6.3) |
| Completeness (%)| 98.5 (96.6) |
| \( <I/\sigma_I> \) | 21.0 (1.5) |
| \( R_{sym} \)   | 0.11 (0.74) |
| \( R_{b_Character}\) | 0.05 (0.32) |
| \( CC_{1/2} \)   | 1.00 (0.80) |

| Refinement statistics |
|-----------------------|
| Resolution (Å)        | 49.52–2.00 (2.02–2.00) |
| Refs used in refinement | 91,238 (2,795) |
| \( R_{work} \) (%)    | 19.9 |
| \( R_{free} \) (%)    | 20.2 |
| Protein atoms         | 6,555 |
| Waters                | 746 |
| Other                 | 0 |

| B value (Å²) |
|--------------|
| Average B value | Protein |
|               | 36 |
|               | Water  |
|               | 43 |
| Wilson B      | 31 |

| RMSD |
|------|
| Bond length (Å) | 0.009 |
| Bond angles (º) | 1.144 |

| Ramachandran plots (%) |
|------------------------|
| Favored                | 97.5 |
| Outliers               | 0.2 |
| PDB                    | 6BTJ |

Abbreviations: APS, Advanced Photon Source; Fab, fragment antigen binding; PDB, Protein Data Bank; RMSD, root-mean-square deviation.

Values in parentheses are for the highest-resolution shell.

\(^a R_{sym} = \Sigma_{n_{id}} \Sigma_i | I_{hkli} - <I_{hkli}> | / \Sigma_{n_{id}} \Sigma_i I_{hkli}\) and

\(^b R_{pim} = \Sigma_{n_{id}} (1/(n-1))^{1/2} \Sigma_i | I_{hkli} - <I_{hkli}> | / \Sigma_{n_{id}} \Sigma_i I_{hkli}, \) where \( I_{hkli} \) is the scaled intensity of the \( i \)th measurement of reflection h, k, l, m, \( <I_{hkli}> \) is the average intensity for that reflection, and \( n \) is the redundancy (1).

\(^c CC_{1/2} = \) Pearson Correlation Coefficient between two random half data sets.

\(^d R_{work} = \Sigma_{n_{id}} | F_o - F_c | / \Sigma_{n_{id}} | F_o | \times 100.\)

\(^e R_{free} = \) was calculated as for \( R_{work} \) but on a test set comprising 5% of the data excluded from refinement.

\(^f \) Calculated using MolProbity (2).

https://doi.org/10.1371/journal.pbio.3000139.t001

(Fig 1B), although the separation of the HA heads make it distinct from the closed, prefusion conformation observed in a large number of crystal structures. This relatively stable complex enabled us to image a larger number of intact particles. In comparisons of 2D classes of cleaved and uncleaved trimeric complexes of H7 HA with H7.5, there appears to be no large difference...
whether zero, one, two, or three Fabs are bound (Fig 1B). Hence, the uncleaved H7 HA, which is stable in the presence of H7.5, is likely to adopt an overall similar structure to the cleaved H7 HA prior to H7.5-induced degradation. We do note that our higher resolution analysis discussed below does indicate that there are likely subtle differences in the molecular interactions between protomers of cleaved and uncleaved HA, depending on the stoichiometry of H7.5 binding.

Our EM data demonstrated that H7.5 had a substantial disruptive effect on the prefusion structure of the HA trimer. To further validate the influence of H7.5 on the HA trimer structure, we investigated the susceptibility of H7 HA to trypsin protease digestion with or without H7.5 bound. Indeed, this experiment showed that addition of H7.5 Fab to H7 Sh2 HA in the presence of trypsin resulted in degradation of HA into many peptidic fragments (S3 Fig). When H7.5 Fab was added to H7 Sh2 HA, 0.1% trypsin was enough to induce the cleavage of H7 Sh2 HA, and 2% was sufficient to completely degrade the H7 HA trimer. Without H7.5 Fab, the H7 Sh2 HA trimers remained resistant to protease cleavage up to a trypsin concentration of 2%. These data indicate that H7.5 may prematurely trigger structural changes or fluctuations of H7 HA trimers that acquire protease-sensitive conformations, which is reminiscent of the typical behavior of HA trimers in low-pH environments [11,12].

To interrogate the structure of H7.5 bound to H7 HA at a higher resolution, we employed single particle cryoEM of the uncleaved H7 NY HA in complex with H7.5 Fab. Reference-free 2D classification resulted in many different views of the complex, and secondary structure features were clearly visible (Fig 2A). The cryoEM 2D classes of the complex in vitreous ice were similar to those observed in nsEM, although the majority of the classes appeared to have three Fabs bound to the HA trimers. Projection images corresponding to meaningful 2D classes were subjected to iterative angular reconstitution and reconstruction. The resulting density
map exhibited significant heterogeneity in the Fab densities attached to the head as well as in the membrane-proximal part of the stem. 3D classification was then performed, and among the resulting classes, one class was characterized by having well-resolved Fabs bound to the head domain. Data in this class were subjected to further refinement resulting in a 3-fold symmetric map with a global resolution of approximately 3.5 Å (EMDB-9139, PDB 6MLM) that is well resolved in all but the membrane-proximal stem region (Fig 2B, Table 2). An initial

Table 2. EM data collection and refinement statistics.

| Map         | H7 HA0 Stem | H7 NL HA1/2-3 H7.5 Fabs | H7 NL HA1/2-3 H7.5 Fabs | H7 SH HA1/2-3 H7.5 Fabs | H7 NY 1/2-3 H7.5 |
|-------------|-------------|-------------------------|-------------------------|-------------------------|------------------|
| EMDB ID     | EMD-9139    | EMD-9139                | EMD-9143                | EMD-9145                | EMD-9142         |
| Data collection |            |                         |                         |                         |                  |
| Microscope  | FEI Titan Krios | FEI Titan Krios          | FEI Talos Arctica       | FEI Talos Arctica       | FEI Talos Arctica |
| Voltage (kV) | 300         | 300                     | 200                     | 200                     | 200              |
| Detector    | Gatan K2 Summit | Gatan K2 Summit         | Gatan K2 Summit         | Gatan K2 Summit         | Gatan K2 Summit  |
| Recording mode | Counting    | Counting                | Counting                | Counting                | Counting         |
| Magnification (incl. postmagnification) | 49,020    | 49,020                   | 43,478                  | 43,478                  | 43,478           |
| Movie micrograph pixel size (Å) | 1.02       | 1.02                    | 1.15                    | 1.15                    | 1.15             |
| Dose rate (e−/(camera pixel) s) | 10         | 10                      | 8.375                   | 8.375                   | 8.375            |
| Number of frames per movie micrograph | 35         | 35                      | 32                      | 32                      | 32               |
| Frame exposure time (ms) | 200       | 200                     | 250                     | 250                     | 250              |
| Movie micrograph exposure time (s) | 7          | 7                       | 9                       | 9                       | 8                |
| Total dose (e−/Å²) | 67         | 67                      | 65                      | 65                      | 60               |
| Defocus range (µm) | 0.4–4.0   | 0.4–4.0                  | 0.4–5.0                 | 0.4–5.0                 | 0.5–4.0          |
| EM data processing |            |                         |                         |                         |                  |
| Number of movie micrographs | 1,324     | 1,324                    |                         |                         |                  |
| Number of molecular projection images in map | 30,032    | 30,032                   | 25,146                  | 9,682                   | 32,117           |
| Symmetry     | C3          | C3                      | C1                      | C1                      | C1               |
| Map resolution (FSC 0.143; Å) | 3.5        | 3.9                     | 9.2                     | 10.2                    | 7.4              |
| Map sharpening B factor (Å²) | −100     | −157                     | −259                    | −109                    | −296             |
| Structure building and validation |            |                         |                         |                         |                  |
| PDB ID       | 6MLM        |                         |                         |                         |                  |
| Number of atoms in deposited model | 16,710    | 5,040                    |                          |                         |                  |
| HA1          | 2,399       | 517                      |                         |                         |                  |
| HA2          | 1,370       | 1,163                    |                         |                         |                  |
| Glycans      | 42          | 0                        |                         |                         |                  |
| MolProbity score | 1.23 (99%) | 1.66 (91%)               |                         |                         |                  |
| Clashscore   | 2.41        | 4.96                     |                         |                         |                  |
| EMRinger score | 2.71      | 1.14                     |                         |                         |                  |
| Deviations from ideal, RMSD |            |                         |                         |                         |                  |
| Bond length (Å) | 0.013     | 0.015                    |                         |                         |                  |
| Bond angles (°) | 1.17      | 1.4                      |                         |                         |                  |
| Ramachandran plot |          |                         |                         |                         |                  |
| Favored (%)  | 96.7        | 94.0                     |                         |                         |                  |
| Allowed (%)  | 3.0         | 5.0                      |                         |                         |                  |
| Outliers (%) | 0.3         | 1.0                      |                         |                         |                  |

Abbreviations: EM, electron microscopy; EMDB, Electron Microscopy Data Bank; FSC, Fourier shell correlation; HA, hemagglutinin; PDB, Protein Data Bank; RMSD, root-mean-square deviation.

https://doi.org/10.1371/journal.pbio.3000139.t002
homology model was created using an X-ray structure of the H7 HA1/HA2 protomer (A/New York/30732-1/2005, 3M5G) [13], combined with our X-ray structure of the H7.5 Fab, and then individually docked into our cryoEM density map exhibiting a nice fit (Fig 2B, 2C and 2D). Next, iterative rebuilding and refinement was carried out in Rosetta [14], yielding a high-resolution atomic model. Since the membrane-proximal part of the map was not well resolved, a protocol for density-subtracted refinement of this region alone was pursued, resulting in a local density map at approximately 3.9 Å resolution (S4 Fig). The membrane-proximal part of the model was refined under constraints of this density map, and an atomic model was created from combining the two builds. To confirm that local (Brownian) motion was responsible for the disorder in the membrane-proximal region—as opposed to the two maps displaying H7 HA in different conformations—coordinates assigned in the density-subtracted local refinement of cryoEM data were applied to the original data, and a reconstruction was performed. Indeed, density corresponding to HA1 and to the parts of HA2 not included in local refinement were clearly visible, albeit at an expected lower resolution (S5 Fig), thereby justifying combining the two builds into a meaningful global conformation supported by our experimental observation.

The epitope on the H7 HA trimer recognized by H7.5 Fab was delineated based on the cryoEM model (Fig 3A). Intriguingly, the H7.5 antibody was found to simultaneously interact with two separated surfaces on two adjacent HA protomers (Figs 2D and 3B). Moreover, our model illustrated that the epitope recognized by H7.5 is not completely accessible when all three HA1 heads are close together, as observed in the apo cleaved H7 trimer crystal structure. Interestingly, the heavy-chain framework region 3 (H-FR3) of H7.5 was juxtaposed to the RBS of an adjacent HA protomer (S6 Fig), and in a closed-trimer model, CDRH3 of H7.5 would clash with residues 189 to 194 in the 190 helix of the adjacent protomer (Fig 3D).

Compared to the apo cleaved H7 trimer model, the buried interprotomer surfaces in the H7.5 epitope were separated substantially from each other in the H7.5-bound H7 NY HA model (Fig 3B). The recessed nature of this epitope suggests that one of the surfaces was likely the primary binding interface recognized by H7.5 antibodies, while the other surface allosterically alters upon antibody binding. Indeed, further interface analysis revealed that the HA surface interacting with CDRH2 and CDRL3 of H7.5 (surface 1) generates a total buried surface area of 720 Å², with substantial favorable energy estimated to be ~5 kcal/mol and an extensive hydrogen bond network between H7.5 and the HA in surface 1 (Fig 3B and 3C). In contrast, the surface between the adjacent HA protomer and H-FR3 of H7.5 (surface 2) is significantly smaller (only 240 Å²) and appears to be energetically unfavorable with a ΔG (change in Gibbs free energy) of +1.2 kcal/mol (Fig 3B and S6 Fig).

To confirm the structural observations, we performed mutagenesis studies of residues in the antibody paratope by individually reverting the residues to the corresponding germline residues and measuring the affinity changes of the H7.5 mutants to H7 Sh2 HA. Consistent with the interface analysis, mutations in CDRH2 and CDRL3, particularly H-S58W, H-T57A, L-T94Q, and L-Y96A of H7.5, drastically reduced binding to H7, while mutations in H-FR3 of H7.5, namely H-M73R and H-S74K, which are proximal to surface 2, did not impact the binding affinity (Fig 3E and S6 Fig). These results strongly support the hypothesis that surface 1 is the primary binding surface driven by favorable energy changes, while surface 2 is a collateral result of the antibody-binding event.

Although mutations in H-FR3 or CDRL3 of the H7.5 antibody did not affect binding of the antibody itself, Fab binding requires the adjacent protomer to be displaced from the tight 3-fold axis at the apex of the trimer to reveal the full epitope (Fig 3D and S6 Fig). The H7.5 epitope therefore includes residues in the inter-HA head contact region (Fig 3E and S6 Fig). Notably, these residues are conserved across all human and zoonotic H7 strains from 1996–2017.
Fig 3. HA-protoner movement and mutagenesis. Each H7.5 Fab was found to recognize a primary epitope on one HA1 subunit and a second smaller region on an adjacent HA1 subunit. (A) The variable domain of one H7.5 Fab is shown as a purple surface for the heavy chain and as a light purple surface for the light chain. The HA1 subunit primarily recognized by H7.5 Fab is shown as a dark green surface, with the epitope surface color in orange (surface 1). The adjacent subunit that is also contacted by the same Fab was colored in light green, with the epitope region in yellow (surface 2). (B) Top view of the H7.5 epitope on H7 HA before and after H7.5 binding. The estimated buried area surfaces on each HA1 subunit and its predicted Gibbs free energies upon H7.5 binding were analyzed. (C) Interaction between Fab H7.5 and HA primary binding surface 1. The H7.5 antibody is shown as main-chain trace, with the interacting residues that are critical to the H7 recognition shown in sticks. The HA1 subunit with surface 1 is shown in a main-chain trace. The main chain and side chain of H7 HA residues that form direct contacts with H7.5 are shown in stick representation. (D) Interaction between two adjacent protoners and a single Fab. When the H7.5 antibody (purple) is docked onto the closed HA (green) head conformation exhibited in 3M5G, there is a clash with the adjacent protoner (top panel). When the 3M5G HA head is morphed into our model, the increased spacing between the heads alleviates this clash, thereby accommodating the CDRH3 loop (bottom panel). (E) The $K_d$ values of H7.5 or mutated H7.5 Fab binding to the HA1 subunit of H7 (A/Shanghai/2/2013) by biolayer interferometry. (F) Sequence alignment of H7 strains with H3N2 reference strain showing the H7.5 epitope (highlighted in yellow) is well conserved among H7. (G) Amino acid sequence analysis of H7.5 epitope among 13,880 influenza A HA sequences obtained from www.fludb.org showing poor conservation in the H7.5 epitope in subtypes other than H7. CDRH3, complementarity determining region heavy chain 3; Fab, fragment antigen binding; HA, hemagglutinin.

https://doi.org/10.1371/journal.pbio.3000139.g003
but not amongst other strains, with the exception of proline at position 62. (S7 Fig). Binding of all three Fabs ultimately results in a conformation of HA with the heads splayed apart.

To further probe the conformational dynamics of the H7 HA when it interacts with H7.5 Fab, we performed hydrogen–deuterium exchange mass spectrometry (HDX-MS) studies on cleaved or uncleaved H7 NL HA in the presence or absence of H7.5 Fab. The differential HDX-MS results of H7 NL HA and H7 NL HA/H7.5 complex were mapped onto the H7 NL HA crystal structure (PDB 4FQV) (Fig 4A). The HDX-MS data revealed two protected regions (colored in blue) in HA1 of H7 HA upon H7.5 binding. These two discontinuous regions (residues E150–K166 and residues S197–L201) form the potential binding epitope, which is consistent with the cryoEM results (Fig 3C). Moreover, HDX-MS experiments performed on cleaved H7 NL HA alone indicated a low level of exchange in one of the regions (residues E150–K166), suggesting less solvent accessibility in the binding epitope region. On the other hand, H7.5 binding led to deprotection of regions (colored in red) in HA1 and HA2, which indicated that HA became more conformationally flexible to accommodate the binding of multiple H7.5 Fabs.

The structural dynamic differences between cleaved and uncleaved H7 NL HA were also compared (Fig 4B). The HDX-MS results indicated that the cleaved H7 NL HA showed more protection (colored in blue) in the HA2 subunit compared to uncleaved H7 NL HA, which is more prone to protease cleavage. This finding is consistent with the helical domains in cleaved HA being more stable and is likely related to the maturation of HA from uncleaved HA0 to HA1 and HA2. Detailed results are shown in S8 Fig.

To trap additional structural intermediates of HA, we next attempted cryoEM reconstructions of H7.5 Fab bound to cleaved H7 Sh2 HA and H7 NL HA, which differ greatly in season and global location of isolation. Guided by our nsEM experiments (Fig 1B), we succeeded in capturing an intermediate structure of cleaved H7 HA using a short incubation time with H7.5 Fab immediately prior to sample vitrification for EM experiments. The predominant classes that we observed had three H7.5 Fabs bound to H7 HA and were able to generate asymmetric 3D reconstructions at approximately 7.4 and approximately 9.2 Å resolution (EMDB-9142, EMDB-9143) (Fig 5C and 5D, S9 Fig and Table 2). Similar to the high-resolution structure of H7.5 bound to uncleaved H7 HA, we still observed movement of the HA1 heads away from the apical 3-fold axis in the cleaved H7 HAs. Further, in all reconstructions of the cleaved H7 HAs, one of the HA1 heads appeared to be splayed out further than the other two (Fig 5C and 5D). In the case of cleaved H7 NL HA, we also refined a class with only two Fabs bound (S9C Fig).

We next compared our high-resolution cryoEM-derived model of the H7.5 bound to H7 NY HA, with separated heads to a crystal structure of HA in the canonical closed prefusion conformation (PDB 3M5G). Within a single protomer, our hybrid docking revealed that the HA1 movement was coupled to a subtle change in its position relative to HA2 (Fig 5). At the trimer level, head separation was accompanied by movement in the HA2 central helices (Fig 5A). Hence, the HA head motions appear to be communicated downward to the stem, resulting in a small separation of the stem protomers. We hypothesize that this looser packing may trigger events that ultimately liberate the fusion peptide or otherwise destabilize the HA trimer and would lead to initiation of the downhill fusion process, a phenotype that would explain our initial nsEM results with the cleaved HA showing dissociation of the trimer into protomers after incubation with H7.5. Additionally, the head separation correlates with HA2 residues G57–K58 adopting an α-helical conformation (Fig 5B). These two amino acids were observed previously as a random coil following the C-terminus of HA2 helix A [13]. The segment (residues G57–I73) connects helix A with the HA2 central helix; in the stable, trimeric HA2
postfusion state, this segment was observed to be \( \alpha \)-helical, forming a continuous extended helix with helix A and the central helix [15]. The \( \alpha \)-helical organization of residues G57–K58 also correlates with further separation of the membrane-proximal H7 stem. Thus, this structure likely represents an intermediate conformational state of the class I fusion protein of

---

**A. Influence of H7.5 Ab on H7 HA Trimer**

| Cleaved Trimer + H7.5 | Uncleaved Trimer + H7.5 |
|-----------------------|------------------------|
| ![Cleaved Trimer + H7.5](4FQV) | ![Uncleaved Trimer + H7.5](4FQV) |
| H7.5 bound trimer | H7.5 bound trimer |

**B. Difference between Cleaved and Uncleaved H7 HA Trimer**

| ![Cleaved HA](4FQV) | ![Uncleaved HA](4FQV) |
|---------------------|---------------------|
| H7.5 bound trimer | H7.5 bound trimer |

---

**Fig 4. HDX-MS analysis of H7 HA trimer–H7.5 complexes.** (A) Difference in HDX profiles of cleaved (left) and uncleaved (right) H7 HA upon H7.5 binding, mapped onto closed (4FQV) and H7.5-bound (H7.5/H7 NY HA) model of H7 HA. The epitope of H7.5 in the HA head shows greater protection, consistent with antibody binding. H7.5 binding also induces greater exchange in the HA head region interprotomer interface as well as the stem, consistent with structural changes in our models. The effects are stronger in the cleaved HA, consistent with H7.5 induction of trimer dissociation. (B) Regions that are more (blue) or less (red) protected in cleaved H7 HA relative to uncleaved H7 HA are mapped onto closed (4FQV) and H7.5-bound (H7.5/H7 NY HA) model of H7 HA. The stem region of the cleaved HA is more protected from exchange than the uncleaved stem, consistent with maturation of the structure and burying of the fusion peptide in the core of the trimer upon cleavage. HA, hemagglutinin; HDX-MS, hydrogen–deuterium exchange mass spectrometry.

[https://doi.org/10.1371/journal.pbio.3000139.g004](https://doi.org/10.1371/journal.pbio.3000139.g004)
Fig 5. Comparison of H7 strains and movement between unliganded and H7.5 liganded HA trimers. (A) Cross sections of open prefusion trimer (H7.5/H7 NY HA) shows separation in the head domain at N199 (red triangle) and in the stem at K39 (green triangle) when compared to closed prefusion trimer (H7 NY HA) (left side). (B) Residues G57–K58 in HA2 show a transition into a helical conformation after H7.5 binds, as seen previously in the postfusion conformation (PDB 1QU1). Moderate-resolution cryoEM reconstructions of cleaved H7 Sh2 HA1/2 (C) and H7 NL HA1/2 (D). Individual HA1/2 protomers from HA (PDB 3M5G) were fit into the reconstructions to generate pseudoatomic.
influenza virus. Fig 6 summarizes the results and the events leading up to the trimer dissociation when H7.5 is added.

Discussion

With the ever-present risk of new avian influenza outbreaks looming, it is important to explore novel ways to target influenza. The recently identified H7.5 recognizes a new conserved epitope on the HA head that can be targeted by neutralizing antibodies. The H7.5 epitope, conserved throughout H7 HA, resides at the HA head protomer–protomer interface and is only transiently accessible. Neutralizing antibodies that bind the HA1 head often target the RBS, blocking HA from interacting with sialic acid on host cells [16]. Although H7.5 does not bind directly to the RBS, it does however block sialic acid binding to HA [10] and potently blocks intact HA from approaching the cell surface.

The prefusion structure of HA has been described extensively, with HA1 and HA2 tightly interacting in a compact arrangement in both cleaved and uncleaved structures [10,17,18]. Portions of the postfusion HA structure have also been crystallized and described [19,20]. Interestingly, mAb H7.5 induces premature dissociation without exposure to low pH, and the somewhat open conformation of the receptor-binding domains may be reminiscent of early steps in the influenza virus HA-mediated membrane-fusion process. After H7.5 binds, it likely induces progressive opening of the trimer, and as more antibodies bind, the trimer is pushed over the activation barrier required for transition into a postfusion conformation [1]. In fact, there are other recently reported HA head–directed antibodies that have indirectly been shown to induce a similar phenotype [21,22]. Notably, these antibodies do not bind to epitopes that overlap with H7.5 (S10 Fig). Antibody-induced decay of HA, via different head-binding antibodies, likely represents a new mechanism of influenza neutralization inhibition.
To date, HA has been shown to adopt a stable, closed prefusion conformation [17,23]. However, despite significant effort, intermediates in the fusion process have eluded high-resolution characterization. It is likely that these intermediates are highly unstable and rapidly transition to the postfusion form [24]. Altogether, our cryoEM studies reveal that HA appears to be somewhat dynamic in its prefusion state, wherein protomers are able to undergo structural fluctuations. Indeed, we observed movement throughout the HA molecule, including fluctuations in the HA2 stem. The inferred movements of the HA heads to enable antibody H7.5 binding suggest that the HA1 head is “breathing,” and this motion is reminiscent of conformational masking described for the HIV envelope glycoprotein [25–27]. Indeed, a very recent study using single-molecule FRET (Förster resonance energy transfer) observed that H5 HA undergoes reversible conformational changes [28]. The structural fluctuations observed in our study, as well as the smFRET study, indicate that movement of the HA1 heads alters interactions between the HA1/HA2 stem portion and the fusion peptide, which reside in the trimer core. HDX-MS studies of H3 HA liberated from whole virions demonstrated a stepwise progression of acid-induced fusion peptide liberation followed by HA head separation [29]. Whether these movements are similarly recapitulated amongst all strains of HA remains to be investigated, but there are likely to be conserved and coordinated motions, given the critical role that the HA-fusion machine plays in viral entry. Thus, there may be ways to exploit these structural fluctuations and access conserved epitopes that have not previously been appreciated relative to the more canonical RBS and stem antibodies.

Materials and methods

Preparation of recombinant H7.5 antibodies

H7.5 Fab for crystallization was prepared as previously described [30]. In brief, the heavy and light chains of H7.5 were cloned independently into the phCMV3 vector and fused with an N-terminal IgK secretion signal. A His6 tag was added to the C-terminus of the Fab heavy chain. Recombinant cDNAs encoding the Fab heavy and light chains were purified and cotransfected into 293F cells by 293fectin (Invitrogen). After 6–7 days of expression at 37 °C, the Fabs were purified from the supernatant by Ni-NTA Superflow (Qiagen) and monoS chromatography (GE Healthcare).

Crystallization and structure determination of H7.5 Fab

H7.5 Fab was concentrated to 10.0 mg/mL in the buffer of 50 mM NaOAc (pH 5.5) for crystallization screening on our high-throughput robotic CrystalMation system (Rigaku) at The Scripps Research Institute (TSRI) using sitting-drop vapor diffusion. The best crystals grew in the well with 0.1 M MES (pH 6.5, 0.01), M cobalt chloride, and 1.8 M ammonium sulfate as mother liquor at 4 °C. Crystals were cryoprotected with mother liquor supplemented with 15% ethylene glycol. X-ray diffraction data were collected to 2.00 Å resolution on beamline 23-ID-D at the Advanced Photon Source (APS). The diffraction data were processed with HKL2000 [31], and the structure was determined by molecular replacement with initial models of PDB 3N9G and PDB 4KMT in Phaser [32]. Refinements were carried out in PHENIX [33], and model rebuilding was performed manually in Coot [34] and the model validated by MolProbity [35].

Structural analysis

Hydrogen bond and buried molecular surface analyses were calculated using the PDBePISA server of EMBL-EBL. Structure figures were generated by MacPyMOL (DeLano Scientific,
Preparation of H7 HA

Baculovirus-expressed HA was prepared for the study as previously described [3,37,38]. In brief, the HA ectodomain sequence was cloned into the pFastBac vector with an N-terminal gp67 secretion signal peptide, a C-terminal BirA biotinylation site, thrombin cleavage site, foldon trimerization domain, and His$_6$ tag. Recombinant bacmid DNA was generated via the Bac-to-Bac system (Invitrogen), and Baculovirus was generated by transfecting purified bacmid DNA into Sf9 cells. HAs were expressed by infecting the High Five cells with the recombinant virus, shaking at 110 rpm for 72 hours at 28 °C. The secreted HAs were purified from the supernatant via Ni-NTA Superflow (Qiagen) and gel filtration (to collect only the HA trimer). The HA trimer was concentrated in the buffer of 20 mM Tris-HCl (pH 8.0) and 150 mM NaCl and aliquoted for storage.

Trypsin digestion for protease susceptibility

For the protease susceptibility assay, the H7 HA in the experimental group was firstly mixed with H7.5 Fab with molar ratio of 1:3 and diluted to 2 mg/ml (for H7 only) in the buffer of 20 mM Tris-HCl (pH 8.0) and 150 mM NaCl. The control group H7 was diluted with the same buffer to the sample concentration. Both groups of H7 samples were then aliquoted and mixed with 0%, 0.1%, 0.2%, 1%, and 2% of trypsin (dissolved in the same buffer above), and each reaction was incubated at room temperature overnight. The samples were then submitted for sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) analysis to determine the stability of the H7 protein.

Biolayer interferometry

An Octet RED instrument (FortéBio, Inc.) was used to determine $K_d$ of the antibody–antigen interactions by biolayer interferometry. To determine the binding affinity of H7.5 Fab to its ligands, HAs (biotinylated as previously described [3,37]) were immobilized onto streptavidin-coated biosensors (FortéBio, Inc.) and incubated with H7.5 Fab at 62.5–500 nM for 120 seconds for association and then incubated in PBS buffer with 0.2% BSA for dissociate for 120 seconds. The signals for each binding event were measured in real time, and $K_d$ values were calculated by fitting to a 1:1 bivalent analyte model. In this case, the $K_d$ values were estimated to be less than $10^{-3}$ nM, as no dissociation was observed.

nsEM

H7.5 IgG was digested with 4% papain w/w for 4 hours before adding iodoacetamide. The Fab was purified using a protein A column and then added to H7 NY HA with a 3x molar excess of Fab and incubated for 30 minutes at room temperature. The complex was purified using an S200i column (GE Healthcare, Amersham, United Kingdom) and immediately added to 400 mesh copper grids with 2% uranyl formate. Micrographs were taken on a Tecnai Spirit with a TemCam F416 camera using Leginon [39]. Particles were selected using DoG Picker [40] and then stacked and aligned into 2D classes in Appion [41] with MRA/MSA [42]. Particles not representing HA trimers or the complex were removed, and a clean stack was processed in RELION [43].
Cryo-EM
Sample preparation for uncleaved H7 HA was similar to negative stain. Directly after size exclusion chromatography (SEC) purification, the complex was added at 1 mg/mL to 2/2 gold Quantifoil grids with amphipol and frozen in liquid ethane using a Vitrobot. Cryo sample preparation for cleaved H7 HA with H7.5 Fab does not include a column purification step. Complexes were formed, incubated for one minute at room temperature, and immediately frozen.

High-resolution image processing
1,324 micrographs were recorded of the uncleaved H7 NY HA/H7.5 Fab complex on a Gatan K2 summit detector mounted on a Titan Krios operating at 300 kV. Data were collected in counting mode at a nominal magnification of 29,000x. Dose rate was approximately 10 electrons/(camera_pixel’s), and frame exposure time was 200 ms. Total exposure time was 10 seconds, with a total dose of 67 electrons/Å. Movie micrograph frames were aligned using MotionCor2 [44], dose weighted, and integrated. Contrast transfer function (CTF) models were determined in GCTF [45]. Projection image candidates of H7 NY HA/H7.5 were identified using a difference-of-Gaussians picker [40]. The resulting set of candidate projection images were subjected to 2D class averaging implemented in RELION 2.1b1 [43]. 227,202 projection images that generated well-formed class averages were selected for further data processing. Iterative angular reconstitution and reconstruction was attempted. Due to on-symmetry axis preferred orientations, reconstruction artefacts were insurmountable for asymmetric refinement. C3 symmetry was imposed in a second attempt, and a stable, albeit somewhat stretched, 3D reconstruction of the data set was obtained. The data set was then subjected to 3D classification, and a subclass of the data (30,032 projection images) was identified; its reconstruction was characterized by persistent Fab densities—indicating full stoichiometric occupancy—and no obvious reconstruction artifacts. This subclass of data was then refined under C3 symmetry constraint to a final resolution of 3.5 Å. Fab Fv regions were well resolved, as were HA1 and part of HA2. The membrane-proximal stem of HA2 was largely disordered. Attempting to recover a structure of the membrane-proximal stem, a globular mask was placed encompassing the disordered part of the map. Inverting the mask, the ordered part of the map could be isolated from which a mask was created to isolate the constant part of the original map. This constant part map was then used to project onto R2 along projection directions deduced from Euler and X,Y coordinate assignments obtained during the earlier-referenced refinement, followed by subtraction from the original projection images. This density-subtracted data set was subjected to further iterative angular reconstitution and reconstruction, resulting in a density map of the membrane-proximal stem resolved to 3.9 Å. To investigate if the two maps could be combined and interpreted as one instance of multiple only locally diverging—breathing—conformations, the Euler and X,Y coordinates determined in the stem refinement was applied to the corresponding original projection images and the coinciding 3D object reconstructed. This density map exhibited, albeit noisy, density for the constant part of the structure, suggesting that the membrane-proximal part of the stem is breathing locally around the position determined in the local map. Treatment of our cleaved H7 NL HA/H7.5 and H7 Sh2 HA/H7.5 data sets was performed in RELION 2.1b1 and followed a standard procedure, as outlined above, prior to local refinement procedures and as outlined in the supplemental information (S5 Fig). FSC 0.143 between independently refined half sets were 7.4 Å (H7 Sh2 HA/3 H7.5 Fabs), 9.2 Å (H7 NL HA/3 H7.5 Fabs), and 10.2 Å (H7 NL HA/2 H7.5 Fabs).
Model building and refinement

A homology model was created (Modeller; [46]) based on the X-ray structure of HA7 HA1/H2 (A/New York/30732-1/2005; PDB 3M5G) [13]. The model was created by combining with the H7.5 Fab X-ray structure and rigid-body fitted to the cryoEM density map. A fragment library consisting of 200 7mers for each amino acid position in the model was compiled from a nonredundant protein structure database. Iterations of manual and Rosetta fragment-library-based centroid rebuilding-and-refinement was then performed [14]. The resulting model was all-atom-refined under constraints of the density map. Glycans were manually built in COOT [34] and final rounds of real-space refinement performed in PHENIX 1.12 [47]. The membrane-proximal part of HA2 was built similarly in the local map referenced above. The builds were then combined into the final structure. Evaluation of builds were performed in MolProbity [35], EMRinger [48], and Privateer [49] and by the PDB validation server.

HDX-MS

Antigen–antibody complexes were prepared by mixing HA H7 and H7.5 antibodies at 1:1.1 stoichiometric ratio and incubating at room temperature for 30 minutes and then kept at 0 °C. For control experiments, free antigens were also diluted with the same volume of protein buffer and treated the same way as complexes. To initiate HDX reactions, 2 μl of prechilled protein stock solution (free uncleaved HA NL H7, 1.8 mg/ml; H7.5-uncleaved HA NL H7, 4.5 mg/ml; cleaved HA NL H7, 1.6 mg/ml; or H7.5-cleaved HA NL H7, 4.5 mg/ml) was diluted into 4 μl D2O buffer (8.3 mM Tris, 150 mM NaCl, in D2O, pD 7.2) at 0 °C. At the indicated times of 10 seconds, 100 seconds, 1,000 seconds, 10,000 seconds, and 100,000 seconds, the exchange reaction was quenched by the addition of 9 μl of optimized quench solution (6.4 M GuHCl, 1 M TCEP, 0.8% formic acid [pH 2.4]) at 0 °C. After incubating on ice for 5 minutes, the quenched sample was diluted 5-fold with 0.8% formic acid containing 16.6% glycerol, immediately frozen on dry ice, and stored at −80 °C. In addition, undeuterated samples and equilibrium-deuterated control samples were also prepared. All samples were then loaded onto our in-house DXMS apparatus for online digestion and separation. The resulting peptides were directed into an OrbiTrap Elite Mass Spectrometer (Thermo Fisher Scientific, San Jose, California) for DXMS analysis. Instrument settings have been optimized for HDX analysis. The data acquisition was carried out in a data-dependent mode and the 5 or 10 most abundant ions were selected for MS/MS analysis. Proteome Discoverer software was used for peptide identification. The centroids of each peptide was calculated with HDExaminer and then converted to corresponding deuteration levels with corrections for back exchange. The deuteration levels of the peptides were further sublocalized using overlapping peptides by MATLAB program.

Epitope conservation analysis

Nonduplicate HA sequences were fetched from the databases at www.fludb.org in FASTA format. Strain A/Aichi/2/1968 HA gene (Genbank Accession M55059) was then defined as a reference for residue numbering. We then used an in-house script to build a database of every HA sequence’s residues at every position relative to the reference. First, we performed pairwise Clustal alignment of every sequence to the reference sequence [50]. The results were parsed into a database keyed on positions relative to the reference, with gaps notated as subpositions following the last identical residue. We are then able to interrogate the database by providing position numbers and desired residues at the position, providing an output of sequences that meet the search criteria as well as breakdowns of residues at the positions. Analysis of natural HA variants was done, as previously described in Wu and colleagues, 2018 [51]. Briefly, a total
of 6,984 HA sequences made up of a subset of up to 20 sequences per year per subtype were analyzed from the Global Initiative for Sharing Avian Influenza Data (GISAID; https://gisaid.org). Sequences were aligned using MAFFT version 7.157b [52]. Sequence logos were generated by WebLogo [53].

Supporting information

S1 Fig. Potent binding of H7.5 Fab to the targeted H7 HAs. The association and dissociation curves from biolayer interferometry of H7.5 Fab to the targeted H7 HAs Shanghai/2/2013 (A) and Netherlands/219/2003 (B) are presented. $K_d$ values are estimated to be less than $10^{-3}$ nM with 1:1 fitting, as no dissociation of the Fab was observed. As previously reported in Thornburg and colleagues [10], the H7.5 antibody shows small amounts of HAI activity against H7 Eurasian and North American H7 strains (C), as well as neutralizing activity against H7N9 Shanghai but not H3N2 Sydney (D). Fab, fragment antigen binding; HA, hemagglutinin.

S2 Fig. Raw micrographs and 2D class averages of cleaved HAs and complexes with H7.5 in nsEM. Cleaved H7 unliganded trimers in nsEM (top row). H7 Shanghai cleaved in complex H7.5 Fab at 4 °C at different incubation times (middle row). H7 New York and H7 Netherlands in complex with H7.5 Fab for 5 minutes and 3 hours, respectively, at 4 °C (bottom row). Fab, fragment antigen binding; HA, hemagglutinin; nsEM, negative-stain electron microscopy.

S3 Fig. Binding of antibody H7.5 increases the protease susceptibility of H7 HA to trypsin digestion. Purified H7 Sh2 HA trimer was incubated with H7.5 Fab (molar ratio of 1:3) or same volume of buffer. The samples were digested with different percentages of trypsin indicated. SDS-PAGE electrophoresis was used to analyze the H7 Sh2 HA stability. Fab, fragment antigen binding; HA, hemagglutinin.

S4 Fig. Protocol for density-subtracted cryoEM data refinement. (A) After data collection, particle extraction, 2D, and 3D classification, all particles resembling the H7 complex were submitted to 3D refinement, resulting in a reconstruction that had diffuse density corresponding to the stem region at the base of the trimer. We therefore subtracted this region from the map and re-refined particles using focused classification. (B) FSC plots of full map and stem base. (C) Resolution maps of full map and stem base show lower resolution in stem alone. (D) Resolution voxel distribution. (E) Relative angular distribution of full map shows bias in side views. cryoEM, cryo-electron microscopy; FSC, Fourier shell correlation.

S5 Fig. Local cryoEM data refinement of membrane-proximal part of H7 NY HA0 in complex with H7.5 Fab. (A) Standard cryoEM data refinement (iterative regularized likelihood optimization of suitable Bayesian posterior functions) results in Euler angular and x, y translational coordinate assignment ($\Phi, \Theta, \psi, x, y)_i$ for each molecular projection image $i$. (B) As density corresponding to the membrane-proximal part of our H7 NY HA0/H7.5 complex was scattered, we proceeded to subtract density corresponding to the well-ordered part of the complex from each of the projection images $i$. The resulting projection images $(i')$ were then, during subsequent refinement, subjected to coordinate optimization locally around the preassigned coordinates ($\Phi', \Theta', \psi', x', y'$). The resulting density map was well ordered in the membrane-proximal region that was previously disordered. (C) We then applied the local refinement coordinates to the original molecular projection images and performed a 3D
reconstruction ($\Phi', \Theta', \psi', x', y'$). The resulting density map was well ordered in the membrane-proximal region. In addition, density corresponding to the membrane-distal part of H7 NY HA0/H7.5 was recovered, albeit at a lower resolution than observed in the original reconstruction in A. This indicates that the disorder in the membrane-proximal part of the complex observed in the original reconstruction was a result of local positional variation (breathing) due to the open base, wherein stabilizing interaction between adjacent protomers is hampered.

(D) Same 3D reconstructed map as in C but displayed at lower contour level. Density corresponding to both H7.5 Fab and the membrane-distal part of H7 NY HA0 is clearly visible. cryoEM, cryo-electron microscopy; Fab, fragment antigen binding; HA, hemagglutinin.

(TIF)

**S6 Fig.** Surface 2 in the H7.5-interacting epitope that is proximal to H-FR3 of the antibody. H7.5 binds to adjacent protomers. Zoomed view of binding surface 1 (dark gold) on protomer 1, binding surface 2 (light gold), and RBS (green) of protomer 2. Residues shown as sticks on H-FR3 interact with surface 2 and fit into the neighboring RBS. H-FR3, heavy-chain framework region 3; RBS, receptor-binding site.

(TIF)

**S7 Fig.** Sequence logos of H7.5 epitope across different influenza subtypes. Analysis of the targeted epitope of H7.5 antibody from HA sequences, including avian and other zoonotic strains, shows a single conserved proline at 62, except H8, which shows no sequence similarity to H7 in the H7.5 epitope. HA, hemagglutinin.

(TIF)

**S8 Fig.** HDX-MS data. (A) Blue suggests the HA regions that exchange slower upon H7.5 Ab binding, and red suggests regions that exchange faster upon binding. (B) Blue suggests the regions that exchange slower upon H7.5 Ab binding, and red suggests the HA regions that exchange faster upon binding. (C) Blue suggests the HA regions that exchange slower in cleaved HA H7, and red suggests the HA regions that exchange faster in cleaved HA H7. HDX profiles for uncleaved (D) and cleaved (E) HA as a function of time. These analyses indicated regions in the respective forms of HA that are more or less accessible to exchange. The exchange profiles for the regions resolved are overall very similar. HA, hemagglutinin; HDX-MS, hydrogen–deuterium exchange mass spectrometry.

(TIF)

**S9 Fig.** Processing cryoEM data of cleaved HA. (A) Data processing and refinement of H7 Shanghai and H7.5 Fab. (B) Data processing and refinement of H7 Netherlands with three and two H7.5 Fabs bound along with angular distribution. (C) FSC plot of H7 Shanghai and H7 Netherlands bound by two or three H7.5 Fabs. cryoEM, cryo-electron microscopy; Fab, fragment antigen binding; FSC, Fourier shell correlation; HA, hemagglutinin.

(TIF)

**S10 Fig.** Comparison of head-splitting antibodies. Crystal structure of m826 (PDB 5VAG) bound to H1 head was aligned to the H1 head region of our cleaved H7 NY model to compare the epitope with H7.5. There is no overlap between antibodies. PDB, Protein Data Bank.

(TIF)

**Acknowledgments**

We thank Robert Kirchdoerfer for help with data processing and interpretation, Bill Anderson and J.C. Ducom for assistance with microscope management and computational support,
Nicholas Wu for help with sequence alignments, and Lauren Holden for critical reading of the manuscript.

**Author Contributions**

**Conceptualization:** Hannah L. Turner, James E. Crowe, Jr., Ian A. Wilson, Andrew B. Ward.

**Data curation:** Hannah L. Turner, Shanshan Lang, Sandhya Bangaru, Ian A. Wilson, Andrew B. Ward.

**Formal analysis:** Hannah L. Turner, Jesper Pallesen, Shanshan Lang, Andrew B. Ward.

**Funding acquisition:** Ian A. Wilson, Andrew B. Ward.

**Methodology:** Hannah L. Turner, Shanshan Lang, Sandhya Bangaru, Christopher A. Cottrell, Charles A. Bowman, Ian A. Wilson, Andrew B. Ward.

**Project administration:** Ian A. Wilson, Andrew B. Ward.

**Resources:** James E. Crowe, Jr., Andrew B. Ward.

**Software:** Charles A. Crowe, Andrew B. Ward.

**Supervision:** Andrew B. Ward.

**Validation:** Hannah L. Turner, Shanshan Lang, Sarah Urata, Sheng Li, Christopher A. Cottrell, Charles A. Bowman, Ian A. Wilson, Andrew B. Ward.

**Visualization:** Hannah L. Turner, Jesper Pallesen, Sheng Li, Christopher A. Cottrell, Andrew B. Ward.

**Writing – original draft:** Hannah L. Turner, Andrew B. Ward.

**Writing – review & editing:** Hannah L. Turner, Ian A. Wilson, Andrew B. Ward.

**References**

1. Schmidt AG, Therkelsen MD, Stewart S, Kepler TB, Liao H-X, Moody MA, et al. Viral receptor-binding site antibodies with diverse germline origins. Cell. 2015; 161: 1026–1034. https://doi.org/10.1016/j.cell.2015.04.028 PMID: 25959776

2. Lee HS, Noh JY, Song JY, Cheong HJ, Choi WS, Jeong HW, et al. Molecular genetic characteristics of influenza A virus clinically isolated during 2011–2016 influenza seasons in Korea. Influenza Other Respir Viruses. 2018; 12: 497–507. https://doi.org/10.1111/irv.12549

3. Lee PS, Ohshima N, Stanfield RL, Yu W, Iba Y, Okuno Y, et al. Receptor mimicry by antibody F045-092 facilitates universal binding to the H3 subtype of influenza virus. Nat Commun. Nature Publishing Group; 2014; 5: 3614. https://doi.org/10.1038/ncomms4614 PMID: 24717798

4. Henry Dunand CJ, Leon PE, Huang M, Choi A, Chromikova V, Ho JY, et al. Both Neutralizing and Non-Neutralizing Human H7N9 Influenza Vaccine-Induced Monoclonal Antibodies Confer Protection. Cell host & microbe. 2016; 19: 800–813. https://doi.org/10.1016/j.chom.2016.05.014 PMID: 27281570

5. Hong M, Lee PS, Hoffman RMB, Zhu X, Krause JC, Laursen NS, et al. Antibody recognition of the pandemic H1N1 Influenza virus hemagglutinin receptor binding site. J Virol. American Society for Microbiology; 2013; 87: 12471–12480. https://doi.org/10.1128/JVI.01388-13 PMID: 24027321

6. Wu NC, Wilson IA. A Perspective on the Structural and Functional Constraints for Immune Evasion: Insights from Influenza Virus. J Mol Biol. 2017; 429: 2694–2709. https://doi.org/10.1016/j.jmb.2017.06.015 PMID: 28648617

7. Zanin M, Koçer ZA, Poulsen RL, Gabbard JD, Howerton EW, Jones CA, et al. Potential for Low-Pathogenic Avian H7 Influenza A Viruses To Replicate and Cause Disease in a Mammalian Model. Williams BFG, editor. J Virol. American Society for Microbiology Journals; 2017; 91: e01934–16. https://doi.org/10.1128/JVI.01934-16 PMID: 27852855

8. de Vries RP, Peng W, Grant OC, Thompson AJ, Zhu X, Bouwman KM, et al. Three mutations switch H7N9 influenza to human-type receptor specificity. Fernandez-Sesma A, editor. PLoS Pathog.
Potent anti-influenza H7 antibody opens HA receptor-binding head domains

Public Library of Science; 2017; 13: e1006390. https://doi.org/10.1371/journal.ppat.1006390 PMID: 28617868

9. Lam TT-Y, Wang J, Shen Y, Zhou B, Duan L, Cheung C-L, et al. The genesis and source of the H7N9 influenza viruses causing human infections in China. Nature. Nature Publishing Group; 2013; 502: 241–244. https://doi.org/10.1038/nature12515 PMID: 23965623

10. Thornburg NJ, Zhang H, Bangaru S, Sapparapu G, Kose N, Lampey RM, et al. H7N9 influenza virus neutralizing antibodies that possess few somatic mutations. J Clin Invest. American Society for Clinical Investigation; 2016; 126: 1482–1494. https://doi.org/10.1172/JCI85317 PMID: 26950424

11. Ekiert DC, Bhabha G, Elsliger M-A, Friesen RHE, Jongeneelen M, Throsby M, et al. Antibody recognition of a highly conserved influenza virus epitope. Science. American Association for the Advancement of Science; 2009; 324: 246–251. https://doi.org/10.1126/science.1171491 PMID: 19251591

12. Kadam RU, Juraszek J, Brandenburg B, Buyck C, Schepens WBG, Kesteleyn B, et al. Peptide fusion inhibitors of influenza virus. Science. American Association for the Advancement of Science; 2017; 358: 496–502. https://doi.org/10.1126/science.aaan0516 PMID: 28971971

13. Yang H, Chen L-M, Carney PJ, Donis RO, Stevens J. Structures of receptor complexes of a North American H7N2 influenza hemagglutinin with a loop deletion in the receptor binding site. Rey FA, editor. PLoS Pathog. Public Library of Science; 2010; 6: e1001081. https://doi.org/10.1371/journal.ppat.1001081 PMID: 20824086

14. DiMaio F, Song Y, Li X, Brunner MJ, Xu C, Conticello V, et al. Atomic-accuracy models from 4.5-Å cryo-electron microscopy data with density-guided iterative local refinement. Nat Methods. Nature Publishing Group; 2015; 12: 361–365. https://doi.org/10.1038/nmeth.3286

15. Chen J, Skehel JJ, Wiley DC. N- and C-terminal residues combine in the fusion-pH influenza hemagglutinin HA(2) subunit to form an N cap that terminates the triple-stranded coiled coil. Proceedings of the National Academy of Sciences. National Academy of Sciences; 1999; 96: 8967–8972.

16. Stadlbauer D, Nachbagauer R, Meade P, Krammer F. Universal influenza virus vaccines: what can we learn from the human immune response following exposure to H7 subtype viruses? Front Med. Higher Education Press; 2017; 11: 471–479. https://doi.org/10.1007/s11684-017-0602-z PMID: 29159597

17. Wilson IA, Skehel JJ, Wiley DC. Structure of the haemagglutinin membrane glycoprotein of influenza virus at 3 Å resolution. Nature. 1981st ed. 1981; 289: 366–373. PMID: 7464906

18. Skehel JJ, Wiley DC. Receptor binding and membrane fusion in virus entry: the influenza hemagglutinin. Annu Rev Biochem. 2000; 69: 531–569. https://doi.org/10.1146/annurev.biochem.69.1.531 PMID: 10966468

19. Malashkevich VN, Higgins CD, Almo SC, Lai JR. A switch from parallel to antiparallel strand orientation in a coiled-coil X-ray structure via two core hydrophobic mutations. Biopolymers. 2015; 104: 178–185. https://doi.org/10.1002/bip.22631 PMID: 25753192

20. Bullough PA, Hughson FM, Skehel JJ, Wiley DC. Structure of influenza haemagglutinin at the pH of membrane fusion. Nature. Nature Publishing Group; 1994; 371: 37–43. https://doi.org/10.1038/371037a0 PMID: 8072525

21. Lee J, Boutz DR, Chromikova V, Joyce MG, Vollmers C, Leung K, et al. Molecular-level analysis of the serum antibody repertoire in young adults before and after seasonal influenza vaccination. Nat Med. Nature Publishing Group; 2016; 22: 1456–1464. https://doi.org/10.1038/nm.4224 PMID: 27820605

22. Yu F, Song H, Wu Y, Chang SY, Wang L, Li W, et al. A Potent Germline-like Human Monoclonal Antibody Targets a pH-Sensitive Epitope on H7N9 Influenza Hemagglutinin. Cell host & microbe. 2017; 22: 471–483.e5. https://doi.org/10.1016/j.chom.2017.08.011 PMID: 28966056

23. Raymond DD, Bajic G, Ferdman J, Suphaphiphat P, Settembre EC, Moody MA, et al. Conserved epitope on influenza-virus hemagglutinin head defined by a vaccine-induced antibody. Proc Natl Acad Sci USA. National Academy of Sciences; 2018; 115: 168–173. https://doi.org/10.1073/pnas.1715471115 PMID: 29255041

24. Harrison SC. Viral membrane fusion. Virology. 2015; 479–480: 498–507. https://doi.org/10.1016/j.virol.2015.03.043 PMID: 25866377

25. Munro JB, Gorman J, Ma X, Zhou Z, Arthos J, Burton DR, et al. Conformational dynamics of single HIV-1 envelope trimers on the surface of native virions. Science. American Association for the Advancement of Science; 2014; 346: 759–763. https://doi.org/10.1126/science.1254426 PMID: 25298114

26. Ozorowski G, Pallesien J, de Val N, Lyumkis D, Cottrell CA, Torres JL, et al. Open and closed structures reveal allostery and pliability in the HIV-1 envelope spike. Nature. Nature Research; 2017; 15: 690–363. https://doi.org/10.1038/nature23010 PMID: 28700571

27. Acharya P, Lusvarghi S, Bewley CA, Kwong PD. HIV-1 gp120 as a therapeutic target: navigating a moving labyrinth. Expert opinion on therapeutic targets. 2015; 19: 765–783. https://doi.org/10.1517/14728222.2015.1010513 PMID: 25724219
28. Das DK, Govindan R, Nikić-Spiegel I, Krammer F, Lemke EA, Munro JB. Direct Visualization of the Conformational Dynamics of Single Influenza Hemagglutinin Trimers. Cell. 2018; 174: 926–937.e12. https://doi.org/10.1016/j.cell.2018.05.050 PMID: 29961575

29. Garcia NK, Guttmann M, Ebner JL, Lee KK. Dynamic changes during acid-induced activation of influenza hemagglutinin. Structure. 2015; 23: 665–676. https://doi.org/10.1016/j.str.2015.02.006 PMID: 25773144

30. Bangaru S, Zhang H, Gilchuk IM, Voss TG, Irving RP, Gilchuk P, et al. A multifunctional human monoclonal neutralizing antibody that targets a unique conserved epitope on influenza HA. Nat Commun. Nature Publishing Group; 2018; 9: 2669. https://doi.org/10.1038/s41467-018-04704-9 PMID: 29961575

31. Otwinowski Z, Minor W. Processing of X-ray diffraction data collected in oscillation mode. Methods in Enzymology. 1997; 276: 307–326.

32. McCoy AJ, Grosse-Kunstleve RW, Adams PD, Winn MD, Storoni LC, Read RJ. Phaser crystallographic software. J Appl Crystallogr. 2007; 40: 658–674. https://doi.org/10.1101/jcb.20084 PMID: 19461840

33. Adams PD, Afonine PV, Bunkóczi G, Chen VB, Davis IW, Echols N, et al. PHENIX: a comprehensive Python-based system for macromolecular structure solution. Acta Crystallogr D Biol Crystallogr. International Union of Crystallography; 2010; 66: 213–221. https://doi.org/10.1107/S090744909052925 PMID: 20124702

34. Emsley P, Cowtan K. Coot: model-building tools for molecular graphics. Acta Crystallogr D Biol Crystallogr. International Union of Crystallography; 2004; 60: 2126–2132. https://doi.org/10.1107/S090744904019158 PMID: 15572765

35. Chen VB, Arendall WB, Headd JJ, Keedy DA, Immormino RM, Kapral GJ, et al. MolProbity: all-atom structure validation for macromolecular crystallography. Acta Crystallogr D Biol Crystallogr. International Union of Crystallography; 2010; 66: 12–21. https://doi.org/10.1107/S090744909042073 PMID: 20057044

36. Pettersen EF, Goddard TD, Huang CC, Couch GS, Greenblatt DM, Meng EC, et al. UCSF Chimera—a visualization system for exploratory research and analysis. J Comput Chem. 2004; 25: 1605–1612. https://doi.org/10.1002/jcc.20084

37. Lang S, Xie J, Zhu X, Wu NC, Lerner RA, Wilson IA. Antibody 27F3 Broadly Targets Influenza A Group 1 and 2 Hemagglutinins through a Further Variation in VH1-69 Antibody Orientation on the HA Stem. Cell Rep. 2017; 20: 2935–2943. https://doi.org/10.1016/j.celrep.2017.08.084 PMID: 28930686

38. Xu R, Krause JC, McBride R, Paulson JC, Crowe JE, Wilson IA. A recurring motif for antibody recognition of the receptor-binding site of influenza hemagglutinin. Nature Structural & Molecular Biology. Nature Publishing Group; 2013; 20: 363–370. https://doi.org/10.1038/nsmb.2500 PMID: 23396351

39. Suloway C, Pulokas J, Fellmann D, Cheng A, Guerra F, Quispe J, et al. Automated molecular microscopy: the new Legion system. J Struct Biol. 2005; 151: 41–60. https://doi.org/10.1016/j.jsb.2005.03.010 PMID: 15890530

40. Voss NR, Yoshioka CK, Radermacher M, Potter CS, Carragher B. DoG Picker and TiltPicker: software tools to facilitate particle selection in single particle electron microscopy. J Struct Biol. 2009; 166: 205–213. https://doi.org/10.1016/j.jsb.2009.01.004 PMID: 19374019

41. Lander GC, Stagg SM, Voss NR, Cheng A, Fellmann D, Pulokas J, et al. Appion: an integrated, data-base-driven pipeline to facilitate EM image processing. J Struct Biol. 2009; 166: 95–102. https://doi.org/10.1016/j.jsb.2009.01.002 PMID: 19263523

42. van Heel M, Harauz G, Orlova EV, Schmidt R, Schatz M. A new generation of the IMAGIC image processing system. J Struct Biol. 1996; 116: 17–24. https://doi.org/10.1006/jsbi.1996.0004 PMID: 8742718

43. Scheres SHW. RELION: implementation of a Bayesian approach to cryo-EM structure determination. J Struct Biol. 2012; 180: 519–530. https://doi.org/10.1016/j.jsb.2012.09.006 PMID: 23000701

44. Zheng SQ, Palovcak E, Armache J-P, Verba KA, Cheng Y, Agard DA. MotionCor2: anisotropic correction of beam-induced motion for improved cryo-electron microscopy. Nat Methods. Nature Publishing Group; 2017; 14: 331–332. https://doi.org/10.1038/nmeth.4193 PMID: 28250466

45. Zhang K. Gctf: Real-time CTF determination and correction. J Struct Biol. 2016; 193: 1–12. https://doi.org/10.1016/j.jsb.2015.11.003 PMID: 26592709

46. Webb B, Sali A. Comparative Protein Structure Modeling Using MODELLER. Curr Protoc Bioinformatics. Hoboken, NJ, USA: John Wiley & Sons, Inc; 2016; 54: 5.6.1–5.6.37. https://doi.org/10.1002/cpbi.3 PMID: 27322406

47. Adams PD, Grosse-Kunstleve RW, Hung L-W, loerger TR, McCoy AJ, Moriarty NW, et al. PHENIX: building new software for automated crystallographic structure determination. Acta Crystallogr D Biol Crystallogr. 2002; 58: 1948–1954. PMID: 12393927
48. Barad BA, Echols N, Wang RY-R, Cheng Y, DiMaio F, Adams PD, et al. EMRinger: side chain-directed model and map validation for 3D cryo-electron microscopy. Nat Methods. Nature Research; 2015; 12: 943–946. https://doi.org/10.1038/nmeth.3541 PMID: 26280328

49. Agirre J, Iglesias-Fernández J, Rovira C, Davies GJ, Wilson KS, Cowtan KD. Privateer: software for the conformational validation of carbohydrate structures. Nature Structural & Molecular Biology. Nature Publishing Group; 2015; 22: 833–834. https://doi.org/10.1038/nsmb.3115 PMID: 26581513

50. Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, et al. Clustal W and Clustal X version 2.0. Bioinformatics. 2007; 23: 2947–2948. https://doi.org/10.1093/bioinformatics/btm404 PMID: 17846036

51. Wu NC, Yamayoshi S, Ito M, Uraki R, Kawaoka Y, Wilson IA. Recurring and Adaptable Binding Motifs in Broadly Neutralizing Antibodies to Influenza Virus Are Encoded on the D3-9 Segment of the Ig Gene. Cell host & microbe. 2018; 24: 569–578.e4. https://doi.org/10.1016/j.chom.2018.09.010 PMID: 30308159

52. Katoh K, Standley DM. MAFFT multiple sequence alignment software version 7: improvements in performance and usability. Mol Biol Evol. 2013; 30: 772–780. https://doi.org/10.1093/molbev/mst010 PMID: 23329690

53. Crooks GE, Hon G, Chandonia J-M, Brenner SE. WebLogo: a sequence logo generator. Genome Res. Cold Spring Harbor Lab; 2004; 14: 1188–1190. https://doi.org/10.1101/gr.849004 PMID: 15173120