Hemagglutinin Traits Determine Transmission of Avian A/H10N7 Influenza Virus between Mammals

Graphical Abstract



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In Brief

Herfst et al. investigated an outbreak in seals caused by an H10N7 influenza virus. In laboratory experiments, this virus was transmissible via the air between ferrets, similar to human influenza viruses. This was caused by mutations that changed the binding pattern and stability of the influenza virus hemagglutinin surface protein.

Highlights

- Adaptation to seals led to transmission of avian A/H10N7 virus between mammals
- Three substitutions in HA altered receptor-binding preference and changed stability
- Receptor-binding specificity substitutions are located in the 220-loop of the HA
- A/H10N7 mammal transmission requirements resemble those for A/H5N1 and other viruses





Article

Hemagglutinin Traits Determine Transmission of Avian A/H10N7 Influenza Virus between Mammals

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SUMMARY

In 2014, an outbreak of avian A/H10N7 influenza virus occurred among seals along North-European coastal waters, significantly impacting seal populations. Here, we examine the cross-species transmission and mammalian adaptation of this influenza A virus, revealing changes in the hemagglutinin surface protein that increase stability and receptor binding. The seal A/H10N7 virus was aerosol or respiratory droplet transmissible between ferrets. Compared with avian H10 hemagglutinin, seal H10 hemagglutinin showed stronger binding to the human-type sialic acid receptor, with preferential binding to α 2,6-linked sialic acids on long extended branches. In X-ray structures, changes in the 220-loop of the receptor-binding pocket caused similar interactions with human receptor as seen for pandemic strains. Two substitutions made seal H10 hemagglutinin more stable than avian H10 hemagglutinin and similar to human hemagglutinin. Consequently, identification of avian-origin influenza viruses across mammals appears critical to detect influenza A viruses posing a major threat to humans and other mammals.

INTRODUCTION

Influenza A viruses are among the most challenging viruses that threaten both human and animal health. In addition to the diversity of host species, influenza A viruses have a remarkable capacity to evolve and adapt after crossing species barriers, to replicate in and transmit between new hosts. The high frequency of inter-species transmission of influenza A viruses results in a high disease burden in humans, pigs, poultry, and to a lesser extent in other mammalian species (Short et al., 2015).

Avian influenza A viruses are classified in subtypes, based on the antigenic and genetic properties of their hemagglutinin (HA) and neuraminidase (NA) membrane glycoproteins. Currently 16 HA subtypes (H1–H16) and 9 NA subtypes (N1–N9) are known to circulate in wild birds, which are the main reservoir of influenza A viruses (Gao, 2018). Influenza A-like viruses of HL17NL10 and HL18NL11 subtypes, identified in bats, have HA- and NA-like proteins substantially different from their homologs in classical avian influenza A viruses (Ma et al., 2015). Occasionally, influ-

subtypes (N1–N9) are known poultry were the source of their infections (Zhang et al., 2014). The main reservoir of influenza In spring 2014, avian influenza A/H10N7 virus infections

caused an outbreak in harbor seals (*Phoca vitulina*) and gray seals (*Halichoerus grypus*) in North-European coastal waters. Initially, more than 500 dead harbor seals were found off the coasts of western Sweden and eastern Denmark (Krog et al.,

enza A viruses from the wild bird reservoir cause sporadic infec-

In this regard, H10 subtype influenza A viruses from waterfowl

have occasionally infected mammals. An A/H10N4 virus caused

an outbreak in mink in 1984 (Klingeborn et al., 1985), and an A/

H10N5 virus was detected in swine in 2008 (Wang et al., 2012).

A/H10N7 viruses were found to cause sporadic human infections in Eqvpt in 2004 (PAHO, 2004) and in Australia in 2010 (Arzey

et al., 2012). More recently, three human cases of infection

with A/H10N8 were detected between December 2013 and

February 2014 (Chen et al., 2014). These patients, all of whom

were isolated cases, were known to have a history of visiting

live poultry markets before disease onset, suggesting that

tions, or even outbreaks, in new host species.

2015; Zohari et al., 2014), and subsequently the outbreak continued south to seals off the coasts of western Denmark and Germany, resulting in the death of another estimated 1,500–2,000 seals, more than 10% of that seal population (Bodewes et al., 2015). More to the south, in the Netherlands, dozens of seals were found dead toward the end of the outbreak.

Phylogenetic analysis showed amino-acid sequence differences between the HA of seal A/H10N7 viruses and closely related Eurasian avian A/H10N7 viruses and the accumulation of amino-acid substitutions in seal A/H10N7 viruses over time (Bodewes et al., 2016). One of these substitutions was a known mammalian adaptation marker HA Q226L (H3 HA numbering is used throughout the manuscript) in the HA receptor-binding site. Although an avian-origin A/H10N7 virus appeared to have spread efficiently in this new host, neither the transmission route between seals nor the need for adaptation of these avian influenza A viruses to infect and transmit among seals is known. Furthermore, since the known avian-origin A/H10N7 and A/ H10N8 viruses have demonstrated their ability to infect humans, it is possible that the avian-origin mammalian-adapted seal A/ H10N7 virus poses an even greater risk for humans. To address this concern that the A/H10N7 virus could be sufficiently adapted and able to spread between and among other mammals, potentially including humans, the transmissibility of the seal A/H10N7 virus via aerosols or respiratory droplets was investigated in the ferret model. Following transmission of the seal A/H10N7 virus, it was compared with a closely related avian A/H10N7 virus to determine which properties had changed in the transmissible seal A/H10N7 virus. Since the major phenotypic differences were attributed to HA, the three-dimensional structures of the mutant seal H10 HA's and of the complexes that they form with avian- and human-type receptor analogs were determined. Combined with biolayer interferometry and glycan array analyses, the receptor-binding preference of the aerosol or respiratory droplet-transmissible virus for human sialic acid receptor analogs was demonstrated. Data from membrane fusion assays showed that the seal A/H10N7 HA was acid stable, which was recently demonstrated to be of importance for transmission of highly pathogenic avian influenza (HPAI) A/H5N1 and other viruses via the aerosol or respiratory route. We discuss our observations in the light of an avian H10-seal H10 comparison to better understand the importance of particular changes in HA for transmission between mammals.

RESULTS

Transmission of A/H10N7 Virus via Aerosols or Respiratory Droplets between Ferrets

To study the ability of A/H10N7 viruses to transmit via aerosols or respiratory droplets between ferrets, transmission experiments were performed as described previously (Munster et al., 2009). First, the transmissibility of an A/H10N7 virus that was isolated from a lung sample collected from a seal in Germany in October 2014, A/harbor seal/Germany/1/14 (H10sealG) (Bodewes et al., 2015) was studied. Although the consensus sequence of the H10sealG HA showed an avian-type-receptor-binding signature with a glutamine at position 226 and a glycine at position 228, deep sequence analysis showed that variants typical of human-type-receptor-binding sites were present at both positions,



namely Q226L and G228S. In addition, the codon for the residue at position 228 was not found in 13% of the sequence reads of this sample, and this deletion was only present in combination with the Q226L mutation (Bodewes et al., 2015). When H10sealG was tested for transmissibility, one out of four recipient ferrets produced a virus culture-positive throat swab at 5 days post exposure (dpe) (Figures 1 and S1). Serum collected from this animal at 14 dpe had a hemagglutination inhibition (HI) antibody titer of 640, confirming transmission and subsequent replication of H10sealG. When the virus-positive swab from the recipient animal was subjected to sequencing, it turned out that the Q226L substitution was positively selected during transmission via aerosols or respiratory droplets. At the same time, the transmissibility of a recombinant A/harbor seal/Netherlands/PV14-221_TS/2015 (H10sealNL, isolated in January 2015) (Bodewes et al., 2016) was investigated, which already contained a leucine at position 226 and in addition a genetically related avian A/ H10N7 virus that was isolated from a mallard in the Netherlands in 2014, A/mallard/NL/1/2014 (H10mall). In these experiments, the H10sealNL virus was transmitted in 6 out of 8 ferret pairs, significantly more than with H10mall that was transmitted in 1 out of 8 transmission pairs (Fisher's exact test, p = 0.04) (Figures 1 and S1).

HA Is the Main Determinant of Phenotypic Differences

Previously, three phenotypic changes were described that render A/H5N1 influenza viruses transmissible via aerosols or respiratory droplets between ferrets: binding to the humantype a2,6 sialic acid receptors in the upper respiratory tract, HA acid and temperature stability, and high polymerase activity (Imai et al., 2012; Linster et al., 2014). Because of the genetic differences between the genomes of the mallard and seal influenza viruses (Table S1), the contribution of the viral genes to possible phenotypic differences was studied. Minigenome assays in 293T cells, plaque assays in Madin-Darby canine kidney (MDCK) cells and replication curves in seal kidney cells indicated that there was no difference in activity of the polymerase complexes of the seal and mallard viruses (Figure S2). In addition, an enzyme-linked lectin assay (ELLA) demonstrated that there were no apparent differences in NA activity between the mallard and seal influenza viruses (Figure S2). Given that there was no indication of any other gene contribution, further investigations were focused on the HA protein.

Receptor-Binding Specificity and Avidity of A/H10N7 Viruses

During circulation in seals, the avian-origin A/H10N7 viruses acquired mutations that may have facilitated efficient virus spread in the new host and associated transmissibility in the ferret model. To assess the influence of these mutations on the receptor-binding specificity of the A/H10 HA's, the attachment patterns of A/Puerto Rico/8/1934 (PR8) viruses harboring, H10sealNL, H10mall, or mutant H10 HA proteins were characterized using a custom sialoside glycan array composed of diverse $\alpha 2,3$ (#11–79) and $\alpha 2,6$ (#80–135) sialosides, including extended sialosides that have been observed in human and ferret upper airway tissues (Peng et al., 2017). A/Indonesia/5/05 (A/H5N1) and A/Netherlands/213/03 (A/H3N2) HA served as controls for avian and human specificity respectively (Figures 2A and 2B).







Figure 1. Transmission of A/H10N7 Viruses via Aerosols or Respiratory Droplets between Ferrets

(A–C) Transmission experiments are shown for H10sealG (A), H10sealNL (B), and H10mall (C). Four (A) or eight (B and C) donor ferrets were inoculated intranasally with $10^6 \, {\rm TCID}_{50}$ of virus and housed individually in transmission cages. A naive recipient ferret was added to each transmission cage adjacent to a donor ferret at 1 dpi. Virus titers in throat swabs of donors (gray lines) and recipients (black dashed lines) were determined by end-point titration in MDCK cells. Transmission via aerosols or respiratory droplets was observed in 1 out of 4 ferret pairs for H10sealNL, and 1 out of 8 ferrets pairs for H10sealNL, and 1 out of 8 ferret pairs for virus detection.

for binding human-type receptors (Figure 3C), and similar double substitutions were found in the HA's of 1957 and 1968 pandemic viruses by comparison with avian A/H2 and avian A/H3 viruses that were possibly their precursors (Connor et al., 1994; Liu et al., 2009). The

H10sealG Q226L, Δ 228 double mutant had very similar avidity for avian- and human-type receptors (Figure 3D).

BLI measurements on H10mall showed that although it had substantially greater avidity for avian-type receptors than for human-type receptors (Figure 3E), it bound human-type receptors more strongly than H10sealG. A previously isolated A/H10 avian influenza virus (A/mallard/Sweden/51/2002 A/H10N2) showed very similar binding properties to H10mall (Figure 3F). H10sealNL and the H10mall Q226L mutant both had approximately 10-fold higher avidity for human-type receptors than avian-type receptors (Figures 3G and 3H).

The binding properties of the control, human A/H3 A/ Netherlands/213/2003 and the avian A/H5 A/Indonesia/5/2005 were also measured by BLI (Figures 3I and 3J). The 2003 seasonal A/H3 virus only bound human-type receptors, whereas the 2005 avian A/H5 virus only bound avian-type receptors.

Altogether, these results for the HA's of the H10 viruses demonstrated that the Q226L mutation was primarily responsible for the change in receptor-binding specificity of the seal influenza viruses from a preference for $\alpha 2,3$ receptors, to dual specificity for both $\alpha 2,3$ - and $\alpha 2,6$ receptors or a preference for $\alpha 2,6$ receptors.

The Structures of the H10seal and H10mall Mutant HA's and the Complexes that They Form with Avian- and Human-Type Receptor Analogs

Purified seal A/H10N7 HA and its variants were crystallized in 3 different crystal forms under similar conditions and contained either a single or two biological trimers per crystallographic asymmetric unit (Table S2). Three out of the 4 potential glycosylation sites were visible in electron density maps.

The structure of the receptor-binding pocket of the H10sealG HA was similar to that of the human origin A/H10N8 HA (A/

H10mall as well as another avian A/H10 HA (A/mallard/Sweden/ 51/2002 - A/H10N2mall) showed a clear avian-type (a2,3-linked) receptor specificity, although some low binding to a2,6 receptors was observed for H10mall (Figures 2C and 2D). In contrast, H10sealNL showed a mixed specificity for a2,3- and a2,6 receptors, with preferential binding to α2,6 receptors with longer extensions on bi-antennary O-linked glycans (#106-108) and biand tri-antennary N-linked glycans (#115-131) (Figure 2E). To assess the role of the HA Q226L substitution on the receptorbinding preference of A/H10 HA, the effect of this substitution in H10mall was analyzed, and the binding preference of the naturally occurring mutants in the receptor-binding site of H10sealG was also studied. On the glycan array, H10mall with Q226L showed dual binding specificity toward both a2,3- and a2,6 receptors (Figure 2F). H10sealG bound only a2,3 receptors, similar to the two avian A/H10 HA's (Figure 2G), but introduction of Q226L resulted in dual receptor specificity (Figure 2H). This is similar to H10mall with the Q226L substitution and H10sealNL that contained Q226L naturally. When both substitutions Q226L and G228S were introduced in H10sealG, dual receptor specificity was observed, and the avidity for shorter a2,6 targets was increased (Figure 2I). The binding preference of the H10sealG deletion mutant, which lacks an amino acid at position 228 but contains Q226L (H10sealG Q226L,Δ228), was similar to the Q226L mutant (Figure 2J).

Biolayer interferometry (BLI) was used to quantitatively analyze the binding of A/H10 seal influenza viruses, to humanand avian-type receptor analogs. BLI measurements on H10sealG showed that, like avian influenza viruses, it had significantly greater avidity for avian-type receptors than for humantype receptors (Figure 3A). The H10sealG Q226L mutant bound avian- and human-type receptors with similar avidity (Figure 3B). The H10sealG Q226L/G228S double mutant had a preference

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Figure 2. Receptor Specificities of Different Avian, Seal, and Mutant A/H10 HA's

Glycan microarray analysis was used to determine the receptor specificities of A/Indonesia/05/2005 (A-J) (A/H5N1) (A), A/Netherlands/213/2003 (A/ H3N2) (B), A/H10N7 HAmall (C), HA A/H10N2mall (D), H10sealNL (E), H10mall Q226L (F), H10sealG (G), H10sealG Q226L (H), H10sealG Q226L, G228S (I), and H10sealG Q226L,Δ228 (J). The mean signals and standard errors were calculated from 4 independent replicates. The data shown are representative of results from 2 independent assays. α2-3-linked sialosides (glycans 11 to 79 on the x axis) and a2-6-linked sialosides (glycans 80 to 135) are shown. Glycans 1 to 10 are nonsialylated controls. RFU, relative fluorescence units.

type receptor and is associated with high affinity binding (Ha et al., 2001; Xiong et al., 2013a, 2013b). The observed binding mode features hydrophilic Gln-226 forming hydrogen bonds with both the Sia-1-Gal-2 glycosidic oxygen and the 4-hydroxyl group of Gal-2. This correlated with a 1-Å upward movement of the Gln-226 side chain from its position in the unbound HA structure (Figure 4A) as seen before (Xiong et al., 2013a) In addition, the distance between the 130-loop and the 220-loop decreased by about 0.5 Å in the HA-avian-type receptor complex compared with the unbound H10sealG structure. The bound avian-type receptor extends upward and exits the receptorbinding site toward the N terminus of the 190 helix.

In contrast, the HA of H10seaINL, which was transmissible via aerosols or respiratory droplets between ferrets, bound avian-type receptor in a *cis*-configuration similar to the configuration of avian-type receptors bound by the transmissible A/H5 mutant and human A/H7N9 HA (Xiong et al., 2013a, 2013b) (Figure 4B). Compared to the receptor analog bound in the *trans*-configuration, the Sia1-Gal2

Jiangxi-Donghu/346/2013) and to that of H10sealNL (Vachieri et al., 2014) (Figures S3A and S3B). Similarly, the structures of the receptor-binding sites of the H10sealG Q226L mutant and of H10sealG Q226L/G228S double mutant were similar to that of the H10sealNL (Figures S3C and S3D). In contrast, the deletion of residue 228 changed the shape of the 220-loop resulting in a movement of about 2 Å in the position of the Leu-226 C- α (Figure S3E).

Avian-Type-Receptor-Binding Characteristics

The H10sealG has a glutamine at position 226 and binds the avian α 2,3-linked sialic acid receptor in a *trans*-configuration (Figure 4A). This is the hallmark of avian HA binding to avian-

glycosidic bond in the *cis*-configuration is rotated counterclockwise more than 90°. Consequently, instead of extending upward toward the N terminus of the 190 helix in the *trans*-configuration, the avian-type receptor exited from the side of the receptor-binding site in the *cis*-configuration over the 220 loop, almost parallel to the 190 helix (Figure 4B). In the *cis*-configuration the hydrophilic glycosidic bond is accommodated further from the hydrophobic leu-226 residue. The change in the conformation of the bound avian-type receptor from *trans* to *cis* often correlates with our observed decrease in affinity (Xiong et al., 2013a, 2013b)

The avian-type receptor also adopted a *cis*-configuration when bound to the H10sealG Q226L mutant and the H10sealG

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Figure 3. Receptor Binding Properties of Different Avian, Seal, and Mutant A/H10 HAs

Biolayer interferometry data of the binding of A/H10 seal and A/H10 mallard viruses and their variants to human- and avian-type receptor analogs (A-H) and control viruses A/Netherlands/213/2003 (A/H3N2) and A/Indonesia/05/2005 (A/H5N1) (I and J). The symbols shown in each diagram are the experimental data for the fractional saturations of the biosensors with various sugar loading levels by a fixed concentration of virus (1 nM). The solid curve is a fit of the experimental data to the Hill equation, which can be used to calculate an estimate of the binding avidity of a virus for a receptor. The binding of viruses to 30 kDa sialyglycopolymers containing α-2,3-sln (avian-type receptor analog) is in red. The binding of viruses to 30 kDa sialyglycopolymers containing a-2,6-sln (human-type receptor analog) is in blue. The H10sealG Q226L mutant binds avian and human receptors with similar avidity (B). This resulted from a 100-fold decrease in avidity for avian-type receptor and a 100-fold increase in avidity for human-type receptor. The Q226L/G228S double mutant preferred human-type receptor (C), which resulted from a 15,000-fold decrease in avidity for avian-type receptors coupled with a ~10fold increase in avidity for human-type receptor. The Q226L, \Delta228 double mutant has similar avidity for avian and human-type receptors resulting from a 10-fold decrease in avidity for avian-type receptor coupled with 3,000-fold increase in avidity for human-type receptors (D).

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H10sealNL

H10sealG Q226L

Q226L/G228S double mutant as in the complexes formed by H10sealNL (Figures 4C and S4A). Alignments of the receptorbinding sites of the Q226L mutant and the Q226L/G228S mutant compared with that of the H10sealNL HA indicated that the orientation of Sia-1 and Gal-2 in the avian-type receptor complexes was similar.

The H10sealG Q226L, Δ 228 mutant bound avian-type receptor in the *trans*-configuration (Figure S4B). Alignment of the Q226L and Q226L, Δ 228 avian-type receptor complexes indicated that the bound sialic acid occupied a similar position relative to the 130-loop. The 220-loop of the deletion mutant was disordered between residues 217 and 226 (Figure S4B), and only density for sialic acid and galactose could be observed in the complex. The disordered 220 loop was accompanied by repositioning of Leu-226, which may facilitate binding of the receptor in the *trans*-configuration. A similar observation was made with an

Figure 4. Avian-Type-Receptor-Binding Features of H10sealG, H10sealNL, and H10sealG Q226L

(A–C) The conformations of avian-type receptors bound to H10sealG, H10sealNL, and H10sealG Q226L and their schematic representations are shown in (A–C), respectively. The sialic acid, galactose, and N-acetylglucosamine of the receptor analogs are colored in yellow, blue, and red, respectively. The receptor-binding site of the avian-type receptor complex of H10sealG is superposed on the receptor-binding site of the apo structure of H10sealG (the distance between C α of 224N and C α of 136A decreased by about 0.5 A) (A). H10sealG is colored in orange in its complex structure and in yellow in its apo structure. HAsealNL is colored in pink (B). H10sealG Q226L mutant is colored in green (C).

avian A/H7 HA with a deleted 220-loop in which the avian-type receptor also adopted a *trans*-configuration (Yang et al., 2010).

Human-Type-Receptor-Binding Characteristics

In line with previous observations made with an A/H5 aerosol or respiratory droplet-transmissible mutant, the Q226L substitution in the receptor-binding site of A/H10 seal HA's correlated with the loss of a hydrogen bond between the side chain of 226Gln and the carboxylate group of Sia-1 (Xiong et al., 2013a). 226Leu interacted with C6 and C4 of the hydrophobic face of the Gal-2 sugar ring, and there is a rotation of \sim 90° about the Sia1-Gal2 glycosidic bond. The Gal-2 and NAG-3 saccharides also rotated ${\sim}90^{\circ}$ toward the 190 helix, and as a result, the plane of the di-saccharide was perpendicular to the plane of the Sia-1 sugar ring. A folded-back conformation of the human-type receptor was observed in

the H10sealG Q226L mutant, the aerosol or respiratory droplet-transmissible H10sealNL, and the H10sealG Q226L/G228S mutant (Figures 5A, 5B, and S5A). This was consistent with tighter human-type receptor binding and with transmission via aerosol or respiratory droplets, properties shared with pandemic A/H1, A/H2, and A/H3 and aerosol or respiratory droplet-transmissible A/H5 HA's (Eisen et al., 1997; Gamblin et al., 2004; Liu et al., 2009; Xiong et al., 2013a).

Previous work has shown that the human-type receptor is bound by the human A/H7N9 HA in a less vertical orientation than the human-type receptors bound to pandemic A/H1, A/ H2, A/H3 HA's, and aerosol or respiratory droplet-transmissible A/H5 HA. It has been proposed that this orientation of humantype receptor bound to A/H7 HA may be due to the 150-loop that protrudes into the receptor-binding site of HA's of the A/ H7, A/H10, and A/H15 clade (Xiong et al., 2013b). However, in





seal A/H10 HA, which also has an extended 150-loop, the human-type receptor adopted a conformation similar to that seen for this receptor in pandemic A/H1, A/H2, A/H3, and aerosol or respiratory droplet-transmissible A/H5 HA's (Figures S6A–S6E).

The H10sealG Q226L mutant and the H10sealG Q226L/ G228S double mutant bind human-type receptor in a similar way to H10sealNL HA (Figures S6F and S6G). Noticeably, the positions of Sia-1 in the different HA monomers of H10sealG Q226L/G228S mutant (carboxylate group rmsd 1.4–2.4Å) are slightly different in comparison with the sialic acids bound in the H10sealG Q226L mutant (carboxylate group rmsd 0.7Å). This heterogeneity is reflected in weaker binding of the humantype receptor observed in BLI.

The H10sealG binds human-type receptor in *cis*-configuration (Figure S4C), similar to wild-type avian A/H5 and A/H7 HA's, with

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Figure 5. Human-Type Receptor-Binding Features of H10sealG Q226L and H10sealNL The side and front views of conformations of human-type receptors bound to H10sealG Q226L and H10sealNL are shown on the left and right sides of (A and B), respectively. The schematic representations of the conformations of the human-type receptors are shown in (A and B) as well. The sialic acid, galactose, and N-acetylglucosamine of receptor analog are colored in yellow, blue, and red individually. H10sealG Q226L is colored in green in its complex structure (A). HAsealNL is colored in pink (B).

the receptor exiting from the front of the site (Xiong et al., 2013a, 2013b). There was a 5° to 10° upward rotation of the glycosidic bond of sialic acid to galactose in H10sealG compared with human-type receptors bound to avian A/H5 and avian A/H7 HA's.

Binding of human-type receptor by the H10sealG Q226L, \Delta228 double mutant involved a disordered 220-loop (Figure S5B), (rmsd 6.5 Å) and of the conformation of the 220-loop in the H10sealG Q226L Δ 228, double mutant and alteration of the position of Leu-226 provided space for the bound human-type receptor to adopt different configurations. By comparison with human-type receptor bound to the H10sealG HA, Gal2-NAG3 residues of the bound human-type receptor in the H10sealG Q226L, Δ228 mutant complex were rotated $\sim 180^{\circ}$ around the glycosidic bond (Figure S5B). In this arrangement the € amino group of Lys137 potentially makes hydrogen bonds with the 5'O and 4'OH of Gal-2 that could stabilize this conformation.

Structural Flexibility and Calcium Binding of H10 Seal HA's

Despite showing overall similarity to the previously characterized mallard

(A/H10N7) and human (A/H10N8) HA structures (Vachieri et al., 2014), the receptor-binding domains of the seal A/H10 HA's varied in positions and orientations relative to the fusion domains (rmsd up to 4.2 Å). By comparison, these relative orientations and positions in human A/H10 HA were less variable (rmsd ~ 0.9 Å). Structural alignment indicated that the relative positions and orientations of the α -helical structure between residues HA₁104 and HA₁114, the 110-helix, also varied in different subunits of the H10sealG Gln226Leu mutant. Examination of this region revealed a calcium-binding site in which the calcium ion was coordinated by the side chains of Glu114 in the 110-helix and of Glu64 of HA₂ in the same subunit (Figure 6). In addition, the side chain of HA₂ Asn79 of a neighboring subunit and the acetyl carbonyl of the first N-acetyl glucosamine in the carbohy-drate side chain attached to HA₂ Asn82 of the same neighboring



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Figure 6. Calcium Binding to A/H10 Seal HA

The calcium-binding site of A/H10 seal HA is shown on the right side. The green sphere represents the calcium ion and the red sphere represents the water molecule. The side chains of the residues, which coordinate with the calcium are shown and the interactions between them and calcium are represented as yellow dash lines. Sugars are colored in green. A ribbon representation of the crystal structure of H10sealG Q226L's complex with avian-type receptor analog is shown on the left side. HA1 and HA2 from one monomer are colored in blue and red, respectively and another two monomers in the HA trimer are colored in gray. The glycosylation in this structure is colored in green. The sialic acid, galactose, and Nacetylglucosamine of the receptor analog are colored in yellow, blue, and red individually.

subunit, also coordinated the calcium. The calcium coordination had pentagonal bipyramidal geometry that closely resembled the calcium-binding site in an EF hand (Kretsinger and Barry, 1975) (Figure 6). Sequence and structural comparisons of the phylogenetically related A/H7 and A/H15 HA's indicated that the orientation of the side chain of Glu114 in the putative calcium-binding sites changed as a result of the absence of His75 from HA₂ of these 2 subtypes (Tzarum et al., 2017b; Xiong et al., 2013b). As a result, calcium was not bound by these HA's.

The observation that calcium coordinates residues of the receptor-binding domain and residues of HA2 of the same and neighboring HA subunits, suggested that bound calcium may influence HA stability. Thermal denaturation experiments gave results consistent with this suggestion by demonstrating that the presence of 2 mM calcium significantly increased the thermal stability of H10sealG 226Q/228G (Tm from 61°C to 69°C), the H10sealG Q226L mutant (Tm from 61°C to 69°C), the H10sealG Q226L, Δ 228 mutant (Tm from 62°C to 67°C), and the H10sealG Q226L/G228S double mutant (Tm from 63°C to 69°C) (Figure S7). In contrast, 2mM calcium did not affect the stability of HA from A/ Aichi/2/68 H3N2 X-31 strain (Figure S7). The results of tryptophan fluorescence assays of denaturation by urea were also consistent with calcium enhancing the stability of A/H10 HA's at room temperature (Figure S8). Calcium stabilized A/H10 human HA and H10sealG Q226L mutant but had little effect on the stability of A/H7 HA and no effect on the stability of X-31 A/ H3N2 HA or A/H5 HA or on the stability of transmissible H10sealNL HA, (Figure S8). The fluorometric titration data indicate that H10sealG and A/H10 human HA bind calcium with micromolar affinities (Figure S9).

The crystal structures of all these HA's suggest a rationale for the results of the stability assays. Electron density for the coordinated calcium ion was observed in crystal structures of A/H10 human, H10mall, and most A/H10 seal HA's. In contrast, the aerosol- or respiratory-droplet-transmissible H10sealNL and A/ H7 HA's in the same clade show no electron density for calcium in their putative binding sites.

The different calcium-binding characteristics observed between H10sealNL and H10sealG could result from two sequence differences in H10sealNL, namely, serine 212 instead of asparagine and isoleucine 244 instead of threonine in the interface between the receptor-binding subdomains. These substitutions influence the relative orientations of the receptor binding, esterase and fusion subdomains, and the position and orientation of Glu114. As a consequence, the loss of interaction between Glu114 and His75 appears to prevent Ca^{2+} binding by H10sealNL.

Despite the effect of calcium binding on HA stability, the results of proteolysis assays indicate that calcium has no detectable effect on the pH of the conformational change of H10sealG Gln226Leu HA (Figure S10) nor does it influence the thermal stability of H10sealG Gln226Leu HA at pH 6.0 (Figure S10).

Membrane Fusion by A/H10N7 Viruses

Attachment of influenza viruses to sialic acid receptors on the cell surface is followed by internalization into endosomes, where, in the low-pH activated conformation, HA mediates fusion of the virus genome into the cytoplasm (Skehel and Wiley, 2000). Previous work has shown that the HA's of aerosol or respiratory droplet-transmissible influenza viruses are comparatively stable, e.g., in studies on mammalian adaptation of avian A/H5N1 viruses. (Herfst et al., 2012; Imai et al., 2012). In another study on the emergence of the swine-origin pH1N1 influenza virus in humans in 2009, it was shown that the HA stability increased during the evolution of A/H1N1 from precursors in swine, to the early pH1N1 human cases and to the later human isolates (Russier et al., 2016).

The fusion pH of H10sealNL and H10mall was estimated in syncytia formation assays. Fusion by H10mall was activated at pH \leq 5.5 (Figure 7). In contrast, the H10sealNL HA required a pH for fusion of <5.2. To determine the substitutions responsible for changes in pH of fusion, all amino acids that were different between H10mall and H10sealNL (18 in total) were introduced individually in H10mall (Table S3). The Q226L substitution in H10mall destabilized HA to pH <5.7. Two-amino-acid substitutions that had the most stabilizing effect on H10mall; T244I and E74D (HA₂) (Table S3) were introduced in H10mall Q226L. This resulted in a decreased pH of fusion of 5.2, suggesting that three-amino-acid substitutions were sufficient to change the receptor-binding specificity and acid stability of H10mall toward that of H10sealNL (Figure 7). As expected, reverse substitutions





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Figure 7. Analysis of the pH Threshold for Fusion of Wild-Type and Mutant A/H10 HA Proteins

Syncytium formation in Vero-118 cells upon expression of wild-type or mutant HA proteins after exposure to different pH. Visual inspection of the cell cultures for the presence of syncytia was used to determine the pH of fusion. The bars indicate the thresholds of pH at which fusion was detected microscopically. The arrows indicate the stability change compared with the parental (WT) H10mall or H10seal. An avian A/HSN1 (A/Indonesia/5/2005) and human A/H3N2 (A/Netherlands/213/2003) HA were included as controls.

in H10sealNL had the opposite effect: L226Q stabilized the HA (pH 4.9), whereas I244T and D74E (HA₂) reverted the stability of the HA (pH 5.4 for both). The triple mutant L226Q, I244T, and D74E (HA₂) in H10sealNL had the same threshold pH of fusion as HAmall, i.e., pH 5.5 (Figure 7).

DISCUSSION

The outbreak of A/H10N7 influenza in seals in 2014 provided an opportunity to study the natural evolution of an avian influenza virus in a mammalian host. This is of interest since it reveals processes involved in the direct transmission of an avian influenza virus to mammals and then between thousands of individuals within a population, and because infected seals could potentially be novel intermediates in the transmission of avian influenza viruses to other mammals including humans. The exact transmission route between seals is unknown, but it is likely to occur via aerosols or respiratory droplets, most probably while the seals are resting on land. However, direct contact transmission between seals can also not be excluded because seals are highly social, and they interact with each other regularly.

Our results indicate that the receptor-binding properties of the HA glycoprotein of the avian influenza virus changed in the establishment of the infection in seals. The avidities and specificities of the changed HA were detailed by microarray analyses using a large number of sialoside receptor analogs, by quantitative, biolayer interferometry assays of receptor analog binding by virus, and by X-ray crystallography of complexes formed between HA and receptor analogs. Together they showed that the changes required for infection of seals caused a decrease in the avidity of the virus for avian-type receptors and an increase in its preference for human-type receptors. The changes were caused by substitutions in the 220-loop that forms one edge of the receptor-binding pocket, in particular the amino-acid substitution Q226L. This change also occurred in the HA of an aerosol or respiratory droplet-transmissible mutant of the initial avian influenza virus that infected seals; it was found when the initial virus replicated in ferrets, and it occurred in viruses isolated in the late phases of the outbreak presumably as a result of selection during the outbreak. This observation is similar to the human A/H2N2 viruses isolated in the first year of the 1957 pandemic. The first viruses to infect humans contained an avian receptorbinding preference, however, likely as the result of adaptation

to replication and transmission in humans, the viruses mutated rapidly, thereby gaining human receptor-binding preference (Pappas et al., 2010). This suggests that the efficient introduction of a new avian influenza virus in mammals does not necessarily require a human receptor-binding phenotype, but this trait may emerge later during the outbreak as the result of mutation and selection.

A/H10N8 viruses were found to cause serious infections of a small number of humans in China in 2013 (Chen et al., 2014). One of the studies of the HA of this virus indicated that by comparison with an avian influenza virus HA that strongly preferred aviantype receptor, the HA of the virus isolated from humans, bound avian and human-type receptors with similar affinity (Vachieri et al., 2014). Another study, however, concluded that the A/ H10N8 virus isolated from infected humans retained the receptor-binding properties of an avian influenza virus (Zhang et al., 2015). The reasons for these different conclusions remain to be determined but they may be related in part to differences in the binding assays and the HA protein used (baculovirus-expressed HA or HA's in virus particles). The study of Zhang et al. also involved site specific substitutions to introduce the amino-acid substitutions Q226L and G228S. The conclusion from microarray receptor-binding assays was that these substitutions were insufficient to change the preference of the A/H10N8 virus for aviantype receptor (Zhang et al., 2015). The human and seal influenza viruses appear, therefore, to differ in this regard also. As well as substitutions in the 220-loop, additional substitutions were made in the 150-loop in a second study, and triple mutants were found to prefer human-type receptors (Tzarum et al., 2017a).

HA in the A/H7, A/H10, A/H15 clade, uniquely among HA's of the five clades into which influenza A viruses are divided on the basis of genetic, antigenic and structural characteristics, contain insertions of amino acids in the 150-loop (Nobusawa et al., 1991), a structural element in the opposite edge of the receptor-binding site from the 220-loop. As a consequence, the extended loops protrude into the receptor-binding site and possibly interfere with binding of human-type receptor in the folded-back conformation (Russell et al., 2004). From the results with the constructed mutants, it was concluded that substitution of residues with shorter amino-acid side chains in the 150-loop resulted in human-type receptor binding (Tzarum et al., 2017a). Again, these would be different requirements from those observed for the A/H10N7 viruses from seals.

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From our observations on the seal influenza viruses, substitutions in the 150-loop are not selected in aerosol or respiratory droplet transmitted mutants, nor, importantly, during the development of the extensive seal outbreak. On the contrary, the mutation Q226L was selected in both cases and this is consistent with the importance of this mutation in the evolution of the HA of the viruses that caused the A/H2 pandemic of 1957 and the A/H3 pandemic of 1968 (Connor et al., 1994).

Substitutions in HA related to the pH of fusion or thermostability have previously been shown to increase aerosol or respiratory droplet transmission of HPAI A/H5N1 and also the 2009 pandemic A/H1N1 (Imai et al., 2012; Linster et al., 2014; Russier et al., 2017). A set of three substitutions, including Q226L, between H10sealNL and H10mall, was also found to affect A/H10 HA stability. Introduction of leucine at residue 226 as a result of mammalian adaptation during circulation in seals resulted in a less stable HA, as previously observed for A/H5N1 and A/H7N9 (Linster et al., 2014; Schrauwen et al., 2016). This destabilizing effect was compensated by T244I and E74D (HA₂) substitutions, which emerged during circulation in the seal population. It is also possible that the HA stability phenotype as measured in fusion and temperature stability assays may co-vary with other, as yet unknown phenotypes, such as stability of HA in aerosols, resistance to drought, stability in mucus, or resistance to changes in pH in the host environment. It should also be noted that H10mall, despite its relatively unstable HA and dual receptorbinding preference for both $\alpha 2,3$ and $\alpha 2,6$ sialic acid, was capable of transmitting through the air, although very inefficient (in one out of eight donor-recipient pairs). Similar findings were obtained recently for a gull-origin A/H10N7 virus that also had a limited aerosol or respiratory droplet-transmissibility in the ferret model (Guan et al., 2019).

The viral phenotypes that changed as a result of the seal adaptation of the avian A/H10N7 virus resemble those that had to change for the aerosol or respiratory droplet-transmissible A/ H5N1 virus, although for A/H10N7, there was no indication that adaptation of the polymerase complex was required. Despite many amino-acid differences between the avian and seal virus polymerase complexes, no differences were observed in the replication kinetics of the avian and seal viruses in seal cells (Table S1 and Figure S2). For adaptation of avian influenza viruses to humans, viral host factors such as importins and ANP32 have been discovered previously, but the possible role of such or other host factors for seals is yet unknown (Gabriel et al., 2008; Long et al., 2016).

H10 subtype influenza viruses have shown the potential not only to infect several mammalian species, including humans, but also to spread efficiently between and among mammals, allowing adaptation to new mammalian hosts. In this study, the adaptation to seals of an avian-origin A/H10N7 virus was studied throughout a series of outbreaks, infecting thousands of individuals, revealing that the same changes in viral properties were required as previously described for HPAI A/H5N1 (Imai et al., 2012; Linster et al., 2014). In addition, changes in HA stability have now been demonstrated for a second avian influenza virus subtype to be of importance to gain aerosol or respiratory droplet-transmissibility between mammals. Our observations help us to increase our basic understanding of influenza virus transmission and also suggest that the emergence of animal influenza viruses with increased HA stability, in addition to human-type receptor preference and



high polymerase activity, should be monitored in surveillance programs aimed to identify zoonotic threats.

STAR***METHODS**

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SUPPLEMENTAL INFORMATION

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AUTHOR CONTRIBUTIONS

S.H., J.Z., J.C.P., J.J.S., and R.A.M.F. conceived, designed, and supervised the project. S.H., J.Z., M.R., R.M., P.L., T.M.B., M.I.J.S., D.d.M., J.M.v.d.B.,



and M.E.R. designed and conducted experiments. S.R.M., S.J.G., X.X., W.P., R.B., E.v.d.V., and A.D.M.E.O. helped with analysis, interpretation of data, and manuscript reviewing and editing. S.H., J.Z., J.C.P., J.J.S., and R.A.M.F. wrote the original draft of the manuscript. All authors read and approved the final manuscript.

DECLARATION OF INTERESTS

Albert Osterhaus is CSO and co-founder Viroclinics Biosciences and CR2O as well as ad hoc consultant, invited speaker, and SAB member to pharmaceutical companies.

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REFERENCES

Adams, P.D., Afonine, P.V., Bunkóczi, G., Chen, V.B., Davis, I.W., Echols, N., Headd, J.J., Hung, L.W., Kapral, G.J., Grosse-Kunstleve, R.W., et al. (2010). PHENIX: a comprehensive Python-based system for macromolecular structure solution. Acta Crystallogr. D Biol. Crystallogr. 66, 213–221.

Arzey, G.G., Kirkland, P.D., Arzey, K.E., Frost, M., Maywood, P., Conaty, S., Hurt, A.C., Deng, Y.M., Iannello, P., Barr, I., et al. (2012). Influenza virus A (H10N7) in chickens and poultry abattoir workers, Australia. Emerg. Infect. Dis. *18*, 814–816.

Bodewes, R., Bestebroer, T.M., van der Vries, E., Verhagen, J.H., Herfst, S., Koopmans, M.P., Fouchier, R.A., Pfankuche, V.M., Wohlsein, P., Siebert, U., et al. (2015). Avian Influenza A(H10N7) virus-associated mass deaths among harbor seals. Emerg. Infect. Dis. *21*, 720–722.

Bodewes, R., Zohari, S., Krog, J.S., Hall, M.D., Harder, T.C., Bestebroer, T.M., van de Bildt, M.W.G., Spronken, M.I., Larsen, L.E., Siebert, U., et al. (2016). Spatiotemporal analysis of the genetic diversity of seal influenza A(H10N7) virus, Northwestern Europe. J. Virol. *90*, 4269–4277.

Chen, H., Yuan, H., Gao, R., Zhang, J., Wang, D., Xiong, Y., Fan, G., Yang, F., Li, X., Zhou, J., et al. (2014). Clinical and epidemiological characteristics of a fatal case of avian influenza A H10N8 virus infection: a descriptive study. Lancet 383, 714–721.

Chutinimitkul, S., van Riel, D., Munster, V.J., van den Brand, J.M., Rimmelzwaan, G.F., Kuiken, T., Osterhaus, A.D., Fouchier, R.A., and de Wit, E. (2010). In vitro assessment of attachment pattern and replication efficiency of H5N1 influenza A viruses with altered receptor specificity. J. Virol. *84*, 6825–6833.

Collaborative Computational Project, Number 4 (1994). The CCP4 suite: programs for protein crystallography. Acta Crystallogr. D Biol. Crystallogr. *50*, 760–763.

Connor, R.J., Kawaoka, Y., Webster, R.G., and Paulson, J.C. (1994). Receptor specificity in human, avian, and equine H2 and H3 influenza virus isolates. Virology *205*, 17–23.

de Wit, E., Munster, V.J., van Riel, D., Beyer, W.E., Rimmelzwaan, G.F., Kuiken, T., Osterhaus, A.D., and Fouchier, R.A. (2010). Molecular determinants of adaptation of highly pathogenic avian influenza H7N7 viruses to efficient replication in the human host. J. Virol. *84*, 1597–1606.

de Wit, E., Spronken, M.I., Bestebroer, T.M., Rimmelzwaan, G.F., Osterhaus, A.D., and Fouchier, R.A. (2004). Efficient generation and growth of influenza virus A/PR/8/34 from eight cDNA fragments. Virus Res. *103*, 155–161.

Dunn, J.J., Krippl, B., Bernstein, K.E., Westphal, H., and Studier, F.W. (1988). Targeting bacteriophage T7 RNA polymerase to the mammalian cell nucleus. Gene *68*, 259–266.

Eisen, M.B., Sabesan, S., Skehel, J.J., and Wiley, D.C. (1997). Binding of the influenza A virus to cell-surface receptors: structures of five hemagglutinin-sialyloligosaccharide complexes determined by x-ray crystallography. Virology 232, 19–31.

Emsley, P., and Cowtan, K. (2004). Coot: model-building tools for molecular graphics. Acta Crystallogr. D Biol. Crystallogr. *60*, 2126–2132.

Gabriel, G., Herwig, A., and Klenk, H.D. (2008). Interaction of polymerase subunit PB2 and NP with importin alpha1 is a determinant of host range of influenza A virus. PLoS Pathog. *4*, e11.

Gamblin, S.J., Haire, L.F., Russell, R.J., Stevens, D.J., Xiao, B., Ha, Y., Vasisht, N., Steinhauer, D.A., Daniels, R.S., Elliot, A., et al. (2004). The structure and receptor binding properties of the 1918 influenza hemagglutinin. Science *303*, 1838–1842.

Gao, G.F. (2018). From "A"IV to "Z"IKV: attacks from emerging and reemerging pathogens. Cell *172*, 1157–1159.

Guan, M., Hall, J.S., Zhang, X., Dusek, R.J., Olivier, A.K., Liu, L., Li, L., Krauss, S., Danner, A., Li, T., et al. (2019). Aerosol transmission of gull-origin Iceland Subtype H10N7 influenza A virus in ferrets. J. Virol. *93*, e00282–19.

Ha, Y., Stevens, D.J., Skehel, J.J., and Wiley, D.C. (2001). X-ray structures of H5 avian and H9 swine influenza virus hemagglutinins bound to avian and human receptor analogs. Proc. Natl. Acad. Sci. USA 98, 11181–11186.

Herfst, S., Mas, V., Ver, L.S., Wierda, R.J., Osterhaus, A.D., Fouchier, R.A., and Melero, J.A. (2008). Low-pH-induced membrane fusion mediated by human Metapneumovirus F protein is a rare, strain-dependent phenomenon. J. Virol. *82*, 8891–8895.

Herfst, S., Schrauwen, E.J., Linster, M., Chutinimitkul, S., de Wit, E., Munster, V.J., Sorrell, E.M., Bestebroer, T.M., Burke, D.F., Smith, D.J., et al. (2012). Airborne transmission of influenza A/H5N1 virus between ferrets. Science *336*, 1534–1541.

Imai, M., Watanabe, T., Hatta, M., Das, S.C., Ozawa, M., Shinya, K., Zhong, G., Hanson, A., Katsura, H., Watanabe, S., et al. (2012). Experimental adaptation of an influenza H5 HA confers respiratory droplet transmission to a reassortant H5 HA/H1N1 virus in ferrets. Nature 486, 420–428.

Klingeborn, B., Englund, L., Rott, R., Juntti, N., and Rockborn, G. (1985). An avian influenza A virus killing a mammalian species-the mink. Brief report. Arch. Virol. *86*, 347–351.

Kretsinger, R.H., and Barry, C.D. (1975). The predicted structure of the calcium-binding component of troponin. Biochim. Biophys. Acta 405, 40–52.

Krog, J.S., Hansen, M.S., Holm, E., Hjulsager, C.K., Chriél, M., Pedersen, K., Andresen, L.O., Abildstrøm, M., Jensen, T.H., and Larsen, L.E. (2015). Influenza A(H10N7) virus in dead harbor seals, Denmark. Emerg. Infect. Dis. *21*, 684–687.

Lin, Y.P., Xiong, X., Wharton, S.A., Martin, S.R., Coombs, P.J., Vachieri, S.G., Christodoulou, E., Walker, P.A., Liu, J., Skehel, J.J., et al. (2012). Evolution of the receptor binding properties of the influenza A(H3N2) hemagglutinin. Proc. Natl. Acad. Sci. USA *109*, 21474–21479.

Linster, M., van Boheemen, S., de Graaf, M., Schrauwen, E.J.A., Lexmond, P., Mänz, B., Bestebroer, T.M., Baumann, J., van Riel, D., Rimmelzwaan, G.F., et al. (2014). Identification, characterization, and natural selection of mutations driving airborne transmission of A/H5N1 virus. Cell *157*, 329–339.

Liu, J., Stevens, D.J., Haire, L.F., Walker, P.A., Coombs, P.J., Russell, R.J., Gamblin, S.J., and Skehel, J.J. (2009). Structures of receptor complexes formed by hemagglutinins from the Asian Influenza pandemic of 1957. Proc. Natl. Acad. Sci. USA *106*, 17175–17180.

Long, J.S., Giotis, E.S., Moncorgé, O., Frise, R., Mistry, B., James, J., Morisson, M., Iqbal, M., Vignal, A., Skinner, M.A., and Barclay, W.S. (2016). Species difference in ANP32A underlies influenza A virus polymerase host restriction. Nature *529*, 101–104.

Ma, W., García-Sastre, A., and Schwemmle, M. (2015). Expected and unexpected features of the newly discovered bat influenza A-like viruses. PLoS Pathog. *11*, e1004819.

Matrosovich, M., Matrosovich, T., Garten, W., and Klenk, H.D. (2006). New low-viscosity overlay medium for viral plaque assays. Virol. J. 3, 63.

Munster, V.J., de Wit, E., van den Brand, J.M., Herfst, S., Schrauwen, E.J., Bestebroer, T.M., van de Vijver, D., Boucher, C.A., Koopmans, M., Rimmelzwaan, G.F., et al. (2009). Pathogenesis and transmission of swineorigin 2009 A(H1N1) influenza virus in ferrets. Science *325*, 481–483.

Nobusawa, E., Aoyama, T., Kato, H., Suzuki, Y., Tateno, Y., and Nakajima, K. (1991). Comparison of complete amino acid sequences and receptor-binding



properties among 13 serotypes of hemagglutinins of influenza A viruses. Virology 182, 475-485.

Osterhaus, A.D., Yang, H., Spijkers, H.E., Groen, J., Teppema, J.S., and van Steenis, G. (1985). The isolation and partial characterization of a highly pathogenic herpesvirus from the harbor seal (Phoca vitulina). Arch. Virol. *86*, 239–251.

PAHO (2004). Avian influenza virus A (H10N7) circulating among humans in Egypt. EID Wkly. Update 2. https://www.paho.org/en/documents/avian-influenza-virus-h10n7-circulating-among-humans-egypt-vol-2-no-18-7-may-2004.

Pappas, C., Viswanathan, K., Chandrasekaran, A., Raman, R., Katz, J.M., Sasisekharan, R., and Tumpey, T.M. (2010). Receptor specificity and transmission of H2N2 subtype viruses isolated from the pandemic of 1957. PLoS One 5, e11158.

Peng, W., de Vries, R.P., Grant, O.C., Thompson, A.J., McBride, R., Tsogtbaatar, B., Lee, P.S., Razi, N., Wilson, I.A., Woods, R.J., and Paulson, J.C. (2017). Recent H3N2 viruses have evolved specificity for extended, branched human-type receptors, conferring potential for increased avidity. Cell Host Microbe *21*, 23–34.

Russell, R.J., Gamblin, S.J., Haire, L.F., Stevens, D.J., Xiao, B., Ha, Y., and Skehel, J.J. (2004). H1 and H7 influenza haemagglutinin structures extend a structural classification of haemagglutinin subtypes. Virology *325*, 287–296.

Russier, M., Yang, G., Marinova-Petkova, A., Vogel, P., Kaplan, B.S., Webby, R.J., and Russell, C.J. (2017). H1N1 influenza viruses varying widely in hemagglutinin stability transmit efficiently from swine to swine and to ferrets. PLoS Pathog. *13*, e1006276.

Russier, M., Yang, G., Rehg, J.E., Wong, S.S., Mostafa, H.H., Fabrizio, T.P., Barman, S., Krauss, S., Webster, R.G., Webby, R.J., and Russell, C.J. (2016). Molecular requirements for a pandemic influenza virus: an acid-stable hemagglutinin protein. Proc. Natl. Acad. Sci. USA *113*, 1636–1641.

Schrauwen, E.J., Richard, M., Burke, D.F., Rimmelzwaan, G.F., Herfst, S., and Fouchier, R.A. (2016). Amino acid substitutions that affect receptor binding and stability of the hemagglutinin of influenza A/H7N9 virus. J. Virol. *90*, 3794–3799.

Short, K.R., Richard, M., Verhagen, J.H., van Riel, D., Schrauwen, E.J., van den Brand, J.M., Mänz, B., Bodewes, R., and Herfst, S. (2015). One health, multiple challenges: the inter-species transmission of influenza A virus. One Health *1*, 1–13.

Skehel, J.J., and Schild, G.C. (1971). The polypeptide composition of influenza A viruses. Virology *44*, 396–408.

Skehel, J.J., and Wiley, D.C. (2000). Receptor binding and membrane fusion in virus entry: the influenza hemagglutinin. Annu. Rev. Biochem. 69, 531–569.

Tzarum, N., de Vries, R.P., Peng, W., Thompson, A.J., Bouwman, K.M., McBride, R., Yu, W., Zhu, X., Verheije, M.H., Paulson, J.C., and Wilson, I.A.

(2017a). The 150-loop restricts the host specificity of human H10N8 influenza virus. Cell Rep. *19*, 235–245.

Tzarum, N., McBride, R., Nycholat, C.M., Peng, W., Paulson, J.C., and Wilson, I.A. (2017b). Unique structural features of influenza virus H15 hemagglutinin. J. Virol. *91*, e00046–17.

Vachieri, S.G., Xiong, X., Collins, P.J., Walker, P.A., Martin, S.R., Haire, L.F., Zhang, Y., McCauley, J.W., Gamblin, S.J., and Skehel, J.J. (2014). Receptor binding by H10 influenza viruses. Nature *511*, 475–477.

Wang, N., Zou, W., Yang, Y., Guo, X., Hua, Y., Zhang, Q., Zhao, Z., and Jin, M. (2012). Complete genome sequence of an H10N5 avian influenza virus isolated from pigs in central China. J. Virol. *86*, 13865–13866.

Watanabe, T., Zhong, G., Russell, C.A., Nakajima, N., Hatta, M., Hanson, A., McBride, R., Burke, D.F., Takahashi, K., Fukuyama, S., et al. (2014). Circulating avian influenza viruses closely related to the 1918 virus have pandemic potential. Cell Host Microbe *15*, 692–705.

Westgeest, K.B., Bestebroer, T.M., Spronken, M.I., Gao, J., Couzens, L., Osterhaus, A.D., Eichelberger, M., Fouchier, R.A., and de Graaf, M. (2015). Optimization of an enzyme-linked lectin assay suitable for rapid antigenic characterization of the neuraminidase of human influenza A(H3N2) viruses. J. Virol. Methods *217*, 55–63.

WHO. (2002). WHO Manual on Animal Influenza Diagnosis and Surveillance (World Health Organization). https://apps.who.int/iris/bitstream/handle/ 10665/68026/WHO_CDS_CSR_NCS_2002.5.pdf?sequence=1&isAllowed=y.

Xiong, X., Coombs, P.J., Martin, S.R., Liu, J., Xiao, H., McCauley, J.W., Locher, K., Walker, P.A., Collins, P.J., Kawaoka, Y., et al. (2013a). Receptor binding by a ferret-transmissible H5 avian influenza virus. Nature *497*, 392–396.

Xiong, X., Martin, S.R., Haire, L.F., Wharton, S.A., Daniels, R.S., Bennett, M.S., McCauley, J.W., Collins, P.J., Walker, P.A., Skehel, J.J., and Gamblin, S.J. (2013b). Receptor binding by an H7N9 influenza virus from humans. Nature *499*, 496–499.

Yang, H., Chen, L.M., Carney, P.J., Donis, R.O., and Stevens, J. (2010). Structures of receptor complexes of a North American H7N2 influenza hemagglutinin with a loop deletion in the receptor binding site. PLoS Pathog. *6*, e1001081.

Zhang, H., de Vries, R.P., Tzarum, N., Zhu, X., Yu, W., McBride, R., Paulson, J.C., and Wilson, I.A. (2015). A human-infecting H10N8 influenza virus retains a strong preference for avian-type receptors. Cell Host Microbe *17*, 377–384.

Zhang, T., Bi, Y., Tian, H., Li, X., Liu, D., Wu, Y., Jin, T., Wang, Y., Chen, Q., Chen, Z., et al. (2014). Human infection with influenza virus A(H10N8) from live poultry markets, China, 2014. Emerg. Infect. Dis. *20*, 2076–2079.

Zohari, S., Neimanis, A., Härkönen, T., Moraeus, C., and Valarcher, J.F. (2014). Avian influenza A(H10N7) virus involvement in mass mortality of harbour seals (Phoca vitulina) in Sweden, March through October 2014. EURO Surveill. *19*, 20967.



STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
NP monoclonal antibody (lgG2a, clone Hb65)	ATCC	Cat# H16L104R5
goat-anti-mouse Ig FITC	BD biosciences, USA	Cat# 349031
Bacterial and Virus Strains		
Influenza virus: H10N7 A/harbour seal/NL/ PV14-221_TS/2015	Bodewes et al., 2016	N/A
Influenza virus: H10N7 A/harbor seal/ S1047_14_L/Germany/2014	Bodewes et al., 2016	N/A
Influenza virus: H10N7 A/mallard/NL/ 1/2014	This paper	N/A
Influenza virus: H10N7 A/Mallard/Sweden/ 51/2002	This paper	N/A
Influenza virus: H10N7 A/harbor seal/ S1047_14_L/Germany/2014 Q226L mutant	This paper	N/A
Influenza virus: H10N7 A/harbor seal/ S1047_14_L/Germany/2014 Q226L, G228S mutant	This paper	N/A
Influenza virus: H10N7 A/harbor seal/ S1047_14_L/Germany/2014 Q226L,∆228 mutant	This paper	N/A
Influenza virus: H10N7 A/mallard/NL/1/ 2014 Q226L mutant	This paper	N/A
Biological Samples		
•		
Turkey red blood cells	In house	N/A
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins	In house	N/A
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS	In house Lonza	N/A Cat# BESP069F
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin	In house Lonza Lonza	N/A Cat# BESP069F Cat# 17-603E
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM)	In house Lonza Lonza Lonza	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate	In house Lonza Lonza Lonza Lonza Lonza	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES	In house Lonza Lonza Lonza Lonza Lonza Lonza	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-737E
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x)	In house Lonza Lonza Lonza Lonza Lonza Lonza Lonza	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-737E Cat#: 13-114E
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin	In house Lonza Lonza Lonza Lonza Lonza Lonza Lonza Sigma	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-737E Cat#: 13-114E T1426-1G
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin Vibrio cholerae neuraminidase (VCNA)	In house Lonza Lonza Lonza Lonza Lonza Lonza Sigma In house	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-737E Cat#: 13-114E T1426-1G N/A
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin <i>Vibrio cholerae</i> neuraminidase (VCNA) Beta-propiolactone	In house Lonza Lonza Lonza Lonza Lonza Lonza Sigma In house Sigma-Aldrich	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-737E Cat#: 13-114E T1426-1G N/A Cat# P5648
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin <i>Vibrio cholerae</i> neuraminidase (VCNA) Beta-propiolactone 2x EMEM	In house Lonza Lonza Lonza Lonza Lonza Lonza Lonza Sigma In house Sigma-Aldrich Lonza	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-737E Cat#: 13-114E T1426-1G N/A Cat# P5648 Cat#:BE12-668F
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin Vibrio cholerae neuraminidase (VCNA) Beta-propiolactone 2x EMEM Avicel RC-591	In house Lonza Lonza Lonza Lonza Lonza Lonza Lonza Sigma In house Sigma-Aldrich Lonza IMCD Benelux BV	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-737E Cat#: 13-114E T1426-1G N/A Cat# P5648 Cat#:BE12-668F RC-591
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin <i>Vibrio cholerae</i> neuraminidase (VCNA) Beta-propiolactone 2x EMEM Avicel RC-591 Phosphate-buffered saline (PBS)	In house Lonza Lonza Lonza Lonza Lonza Lonza Lonza Sigma In house Sigma-Aldrich Lonza IMCD Benelux BV Oxoid	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-737E Cat#: 13-114E T1426-1G N/A Cat# P5648 Cat#:BE12-668F RC-591 X6571D
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin <i>Vibrio cholerae</i> neuraminidase (VCNA) Beta-propiolactone 2x EMEM Avicel RC-591 Phosphate-buffered saline (PBS) OPD substrate	In house Lonza Lonza Lonza Lonza Lonza Lonza Lonza Sigma In house Sigma-Aldrich Lonza IMCD Benelux BV Oxoid Sigma-Aldrich	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-737E Cat#: 13-114E T1426-1G N/A Cat# P5648 Cat#:BE12-668F RC-591 X6571D P8287-50TAB
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin <i>Vibrio cholerae</i> neuraminidase (VCNA) Beta-propiolactone 2x EMEM Avicel RC-591 Phosphate-buffered saline (PBS) OPD substrate fetuin	In house Lonza Sigma In house Sigma-Aldrich Lonza IMCD Benelux BV Oxoid Sigma-Aldrich Sigma-Aldrich Sigma-Aldrich	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-737E Cat#: 13-114E T1426-1G N/A Cat# P5648 Cat#:BE12-668F RC-591 X6571D P8287-50TAB F3385-1G
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin <i>Vibrio cholerae</i> neuraminidase (VCNA) Beta-propiolactone 2x EMEM Avicel RC-591 Phosphate-buffered saline (PBS) OPD substrate fetuin Nunc-Immuno™ MicroWell™ 96 well solid plates	In house Lonza Sigma In house Sigma-Aldrich Lonza IMCD Benelux BV Oxoid Sigma-Aldrich Sigma-Aldrich Sigma-Aldrich Sanbio, Uden, The Netherlands	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-613E Cat#: 17-737E Cat#: 13-114E T1426-1G N/A Cat# P5648 Cat#:BE12-668F RC-591 X6571D P8287-50TAB F3385-1G 442404
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin <i>Vibrio cholerae</i> neuraminidase (VCNA) Beta-propiolactone 2x EMEM Avicel RC-591 Phosphate-buffered saline (PBS) OPD substrate fetuin Nunc-Immuno [™] MicroWell [™] 96 well solid plates PNA-HRPO	In house Lonza In house Sigma-Aldrich	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-737E Cat#: 13-114E T1426-1G N/A Cat# P5648 Cat#:BE12-668F RC-591 X6571D P8287-50TAB F3385-1G 442404 L7759-1MG
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin <i>Vibrio cholerae</i> neuraminidase (VCNA) Beta-propiolactone 2x EMEM Avicel RC-591 Phosphate-buffered saline (PBS) OPD substrate fetuin Nunc-Immuno [™] MicroWell [™] 96 well solid plates PNA-HRPO Xtremegene transfection reagent	In house Lonza Inhouse Sigma-Aldrich Sigma-Aldrich Sigma-Aldrich Sigma-Aldrich Sigma-Aldrich Sigma-Aldrich Roche	N/A Cat# BESP069F Cat# 17-603E Cat#: 17-737E Cat#: 17-737E Cat#: 13-114E T1426-1G N/A Cat# P5648 Cat#:BE12-668F RC-591 X6571D P8287-50TAB F3385-1G 442404 L7759-1MG 06366546001
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Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin <i>Vibrio cholerae</i> neuraminidase (VCNA) Beta-propiolactone 2x EMEM Avicel RC-591 Phosphate-buffered saline (PBS) OPD substrate fetuin Nunc-Immuno™ MicroWell™ 96 well solid plates PNA-HRPO Xtremegene transfection reagent Giemsa 0.05% Trypsin-EDTA (1x)	In house Lonza Sigma In house Sigma-Aldrich Roche Merck Millipore Gibco	N/A Cat# BESP069F Cat# 17-603E Cat# 17-613E Cat# 17-737E Cat# 13-114E T1426-1G N/A Cat# P5648 Cat#:BE12-668F RC-591 X6571D P8287-50TAB F3385-1G 442404 L7759-1MG 06366546001 1092040500 Cat# 25300-054
Turkey red blood cells Chemicals, Peptides, and Recombinant Proteins EMEM with EBBS Penicillin-Streptomycin L-Glutamine (200 mM) sodium bicarbonate 1M HEPES Non-essential amino acids (100x) TPCK-treated trypsin Vibrio cholerae neuraminidase (VCNA) Beta-propiolactone 2x EMEM Avicel RC-591 Phosphate-buffered saline (PBS) OPD substrate fetuin Nunc-Immuno™ MicroWell™ 96 well solid plates PNA-HRPO Xtremegene transfection reagent Giemsa 0.05% Trypsin-EDTA (1x) 6'SLN	In house Lonza Sigma In house Sigma-Aldrich Lonza IMCD Benelux BV Oxoid Sigma-Aldrich Sigma-Aldrich Sigma-Aldrich Sigma-Aldrich Sigma-Aldrich Sigma-Aldrich Roche Merck Millipore Gibco Dextra Laboratories Ltd., Reading, UK	N/A Cat# BESP069F Cat# 17-603E Cat# 17-605F Cat#: 17-613E Cat#: 17-613E Cat#: 17-613E Cat#: 17-737E Cat#: 13-114E T1426-1G N/A Cat# P5648 Cat#:BE12-668F RC-591 X6571D P8287-50TAB F3385-1G 442404 L7759-1MG 06366546001 1092040500 Cat# 25300-054 Cat#SLN306

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Article



Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
3'SLN	Dextra Laboratories Ltd., Reading, UK	Cat#SLN302
6'SLN-PAA	Lectinity Holdings, Moscow, Russia	Cat#0997-BP
3'SLN-PAA	Lectinity Holdings, Moscow, Russia	Cat#0036-BP
Oseltamivir	Kind gift from Roche, Welwyn Garden City, UK	N/A
Zanamivir	Kind gift from GSK, Stevenage, UK	N/A
Trypsin	Sigma	Cat#T1426
H10sealG HA	This paper	N/A
H10sealG Q226L HA	This paper	N/A
H10sealG Q226L,G228S HA	This paper	N/A
H10sealG Q226L, Δ228 HA	This paper	N/A
H10sealNL HA	This paper	N/A
Critical Commercial Assays		
QuikChange multi-site-directed mutagenesis kit	Agilent	Cat# 200514
Dual-Glo Luciferase Assay System	Promega	Cat# E2920
Deposited Data		
H10sealG Q226L, ∆228 HA (apo) structure	This paper	PDB: 6TJW
The structure of H10sealG Q226L, Δ228 HA-3'SLN complex	This paper	PDB: 6TXO
The structure of H10sealG Q226L, Δ228 HA-6'SLN complex	This paper	PDB: 6TVT
H10sealG Q226L HA (apo) structure	This paper	PDB: 6TVR
The structure of H10sealG Q226L HA- 3'SLN complex	This paper	PDB: 6TVS
The structure of H10sealG Q226L HA- 6'SLN complex	This paper	PDB: 6TWV
H10sealNL HA (apo) structure	This paper	PDB: 6TVC
The structure of H10sealNL HA-3'SLN complex	This paper	PDB: 6TVA
The structure of H10sealNL HA-6'SLN complex	This paper	PDB: 6TVB
H10sealG Q226L, G228S HA (apo) structure	This paper	PDB: 6TWH
The structure of H10sealG Q226L, G228S HA-3'SLN complex	This paper	PDB: 6TWI
The structure of H10sealG Q226L, G228S HA-6'SLN complex	This paper	PDB: 6TY1
The structure of H10sealG Q226L, G228S HA- EDTA complex	This paper	PDB: 6TWS
H10sealG HA (apo) structure	This paper	PDB: 6TJY
The structure of H10sealG HA-3'SLN complex	This paper	PDB: 6TVD
Experimental Models: Cell Lines		
Human: HEK 293T cells	N/A	N/A
Canine: MDCK II cells	ATCC	Cat# CCL-34
Seal (Phoca Vitulina) kidney cells	Osterhaus et al., 1985	N/A
Embryonated chicken eggs	Drost	N/A
Spodoptera frugiperda: Cell line Sf9	Thermo Fisher Scientific	Cat#12659017
Experimental Models: Organisms/Strains		
Ferret (Mustela putorius furo)	Euroferret	N/A

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Article

Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Recombinant DNA		
Plasmid: pHW2000 (modified version)	(Chutinimitkul et al., 2010)	N/A
Plasmid: pFB-H10sealG HA	Thermo Scientific	N/A
Plasmid: pFB-H10sealG Q226L HA	Thermo Scientific	N/A
Plasmid: pFB-H10sealG Q226L, ∆228 HA	Thermo Scientific	N/A
Plasmid: pFB-H10sealG Q226L, G228S HA	Thermo Scientific	N/A
Plasmid: pFB-H10sealNL HA	Thermo Scientific	N/A
Software and Algorithms		
ImageQuant TL Software	GE Healthcare Life Sciences	N/A
iMOSFLM		N/A
Aimless		N/A
Pointless		N/A
The CCP4 suite	Collaborative Computational Project, 1994	N/A
Phenix	Adams et al., 2010	N/A
Other		
Minigenome assay	de Wit et al., 2010	N/A
Sialoside glycan microarray	This paper and (Peng et al., 2017)	N/A

RESOURCE AVAILABILITY

Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Ron Fouchier (r.fouchier@erasmusmc.nl).

Materials availability

All unique reagents generated in this study are available from the Lead Contact with a completed Materials Transfer Agreement.

Data and code availability

All datasets for glycan microarray experiments are included in the supplement according to the MIRAGE consortium format (Table S4, supplementary glycan microarray document). These datasets have not been uploaded to a public repository since, at present, such a resource does not exist. Structural data have been deposited with the Protein Data Bank under accession codes 6TJW, 6TXO, 6TVT, 6TVR, 6TVS, 6TWV, 6TVC, 6TVA, 6TVB, 6TWH, 6TWI, 6TY1, 6TWS, 6TJY, 6TVD, 6TVF.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Experimental Animals

Forty female ferrets (*Mustela putorius furo*), 12- to 24- month old, sero-negative for prototype A/H1, A/H3 and A/H5 influenza A viruses and Aleutian Disease Virus, were obtained from an accredited ferret breeder. All animals were microchipped and received hormonal treatment to prevent estrus. Ferrets were housed with a 12 h light/dark cycle and allowed access to diet and water. The experimental set-up was specifically designed to allow transmission experiments to be conducted in negatively pressurized isolator cages (1.6m x 1m x 1m) in the ABSL3+ facility. During the experiment the ferrets are housed in clear Perspex cages, in which each inoculated animal was housed individually next to a naive ferret. Each ferret cage was 30 cm x 30 cm x 55 cm (W x H x L) and the two cages were separated by two stainless steel grids (1), with a grid size of 0.5 cm², 10 cm apart. Experiments were performed in strict compliance with European guidelines (EU Directive on Animal Testing 86/609/EEC) and Dutch legislation (Experiments on Animals Act, 1997). An independent animal experimentation ethical review committee (Dutch Stichting Dier Experimenten Commissie Consult) approved all animal studies.

METHOD DETAILS

Biocontainment

All experiments involving the seal A/H10N7 viruses were conducted at enhanced animal biosafety level 3 (ABSL3+). The ABSL3+ facility of Erasmus MC consists of a negative pressurized (-30 Pa) laboratory in which all *in vivo* and *in vitro* experimental work is



carried out in class 3 isolators or class 3 biosafety cabinets, which are also negative pressurized (< -200 Pa). Although the laboratory is considered 'clean' because all experiments are conducted in closed class 3 cabinets and isolators, special personal protective equipment, including laboratory suits, gloves and FFP3 facemasks is used. Air released from the class 3 units is filtered by High Efficiency Particulate Air (HEPA) filters and then leaves the facility via a second set of HEPA filters. Only authorized personnel that received the appropriate training can access the ABSL3+ facility. All personnel working in the facility is vaccinated against seasonal and A/H5N1 influenza viruses. For animal handling in the facilities, personnel always work in pairs. The facility is secured by procedures recognized as appropriate by the institutional biosafety officers and facility management at Erasmus MC and Dutch and United States government inspectors. Antiviral drugs (oseltamivir and zanamivir) and personnel isolation facilities are directly available to further mitigate risks upon incidents. All experiments described in this manuscript were performed with naturally occurring viruses, and therefor this research does not fall under the pause on gain-of-function research.

Viruses

Influenza viruses A/harbour seal/NL/PV14-221_TS/2015 (H10sealNL) and A/harbor seal/S1047_14_L/Germany/2014 (H10sealG) were propagated in MDCK cells (Bodewes et al., 2016). Influenza virus A/mallard/NL/1/2014 (H10mall) was propagated in embryonated chicken eggs followed by one passage in MDCK cells. All eight gene segments were amplified by reverse transcription polymerase chain reaction and cloned in a modified version of the bidirectional reverse genetics plasmid pHW2000 (Chutinimitkul et al., 2010; de Wit et al., 2004). Substitutions of interest were introduced by reverse genetics using the QuikChange multi-site-directed mutagenesis kit (Agilent Netherlands, Amstelveen, Netherlands) according to the instructions of the manufacturer. Recombinant viruses were produced upon transfection of 293T cells and virus stocks were propagated in MDCK cells or embryonated chicken eggs (Drost, Loosdrecht, The Netherlands) and titrated in MDCK cells. For binding assays and stability assays, reassortant viruses consisting of seven gene segments of influenza virus A/PR/8/34 and the HA segment of interest were produced using a previously described reverse genetics system for influenza virus A/PR/8/34 (de Wit et al., 2004).

Virus Titration in MDCK Cells

Virus titers were determined by end-point titration in MDCK cells. MDCK cells were inoculated with tenfold serial dilutions of virus stocks, nose swabs, or throat swabs. Cells were washed with PBS one hour after inoculation and cultured in 200µl of infection media, consisting of Eagle's Minimum Essential Medium (EMEM) supplemented with 100 U/ml penicillin, 100 µg/ml streptomycin, 2 mM glutamine, 1.5 mg/ml sodium bicarbonate, 10 mM HEPES (4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid), non-essential amino acids, and 20 µg/ml N-tosyl-L-phenylalanine chloromethyl ketone (TPCK) treated trypsin (Sigma). Three days after inoculation, supernatants of infected cell cultures were tested for agglutinating activity using turkey red blood cells (TRBCs) as an indicator of virus replication in the cells. Infectious virus titers were calculated from 4 replicates (nose swabs, and throat swabs) or 10 replicates (virus stocks) by the method of Spearman-Karber.

Transmission of A/H10N7 Virus via Aerosols or Respiratory Droplets in the Ferret Model

We chose the ferret (*Mustela putorius furo*) as the animal model for our studies. Ferrets have been used in influenza research since 1933 because they are susceptible to infection with human and avian influenza viruses. After infection with human influenza A virus, ferrets develop respiratory disease and lung pathology similar to that observed in humans. Ferrets can also transmit human influenza viruses via the air to other ferrets that serve as sentinels with or without direct contact. Aerosol or respiratory droplet transmission experiments were performed as described previously (Herfst et al., 2012; Munster et al., 2009). In short, seronegative female adult ferrets were inoculated intransally with 10^6 50% Tissue Culture Infectious Dose (TCID₅₀) of virus by applying 250 µl of virus suspension to each nostril. Each ferret was then placed in a transmission cage. One day after inoculation, one naïve ferret was placed opposite to each inoculated ferret. Each transmission pair was housed in a separate transmission cage designed to prevent direct contact between the inoculated and naïve ferrets but allowing airflow from the inoculated to the naïve ferret. Nose and throat swabs were collected on 1, 3, 5, and 7 days post inoculation (dpi) for inoculated ferrets and on 1, 3, 5, 7, and 9 days post exposure (dpe) for the naïve ferrets. Virus titers in swabs were determined by end-point titration in MDCK cells. All animals were humanely euthanized at 14 dpi/dpe.

Serology

The presence of antibodies elicited against the tested viruses were confirmed by hemagglutination inhibition (HI) assay using standard procedures (WHO, 2002). Briefly, ferret antisera were prepared following intranasal inoculation, from blood collected 14 days later. Antisera were pre-treated overnight with receptor destroying enzyme *Vibrio cholerae* neuraminidase (VCNA) at 37°C, and incubated at 56°C for 1h the next day. Two-fold serial dilutions of the antisera, starting at a 1:20 dilution, were mixed with 25 µl of a virus stock containing 4 hemagglutinating units and were incubated at 37°C for 30 minutes. Subsequently, 25 µl 1% TRBCs was added and the mixture was incubated at 4°C for 1h. HI was read and was expressed as the reciprocal value of the highest dilution of the serum that completely inhibited agglutination of virus and erythrocytes.

Purification of HAs

The ecto domains of all A/H10 seal HAs were cloned into a pFastBac1 vector. This vector was edited for HA ectodomain expression, which contained a polyhedrin signal peptide at the N-terminal of HA and a tobacco etch virus cleavage site, a foldon for trimerization



and an octa-histidine tag at the C-terminus of HA. All A/H10 seal HAs were expressed in Sf9 cells and purified by a combination of Co-NTA (Co³⁺ complex with nitrilotriacetic acid) chromatography, trypsin digestion and gel filtration chromatography (Lin et al., 2012).

Crystallization

Purified H10 seal HAs were concentrated to 10 mg/ml for crystallization. H10sealG was crystallized in 2%-4% PEG6000, 0.1M HEPES pH7.0 and 8% PEG6000, 22% ethylene glycerol, 0.1M HEPES pH 7.5 was used as a cryoprotectant for flash freezing in liquid nitrogen. H10sealG Q226L was crystallized in 4% PEG3350, 0.1M HEPES pH 7.5 and 15% PEG3350, 20% ethylene glycol, 0.1M HEPES pH7.5 was applied as a cryoprotectant for flash freezing in liquid nitrogen. H10sealG Q226L, Δ228 mutant crystals were grown in 10% PEG6000, 0.1M bicine pH9.0 and flash freezing in liquid nitrogen with the cryoprotectant 12% PEG6000, 22% ethylene glycol, 0.1 M bicine pH9.0. H10sealG Q226L/G228S crystals were grown in 6% PEG3350, 0.1 M MES pH 6.5 and flash freezing in liquid nitrogen with a cryoprotectant 15% PEG3350, 20% ethylene glycol, 0.1 M bicine pH9.0. H10sealG Q226L/G228S crystals were grown in 6% PEG3350, 0.1 M MES pH 6.5 and flash freezing in liquid nitrogen with a cryoprotectant 15% PEG3350, 20% ethylene glycol, 0.1 M HEPES pH 7.5. H10sealNL crystals were grown in 26%-31% PEG600, 0.1M HEPES pH7.5 or 0.1M tris pH8.0 and flash freezing in liquid nitrogen with a cryoprotectant 33% PEG600, 15% ethylene glycol, 0.1M HEPES pH7.5. Crystals were soaked for 3mins-6mins in 100mM 3'-sialyl-N-acetyllactosamine (SLN) or 6'-SLN in cryobuffer. All the data sets were collected at the Diamond Light Source, Harwell, Oxfordshire, UK. Diffraction datasets were processed using Xia2 DIALS, imosflm and scala (Collaborative Computational Project, 1994). Structures were solved by molecular replacement with Phaser and the coordinate file of A/H10 A/Jiangxi-Donghu/346/2013 HA (Protein Data Bank [PDB] entry 4D00) was used as a searching model (Vachieri et al., 2014). The structures were manually built in coot (Emsley and Cowtan, 2004) and refined by using Refmac5 (Collaborative Computational Project, 1994) and Phenix (Adams et al., 2010).

Virus Preparation for Biolayer Interferometry

All the A/H10N7 seal influenza viruses were inoculated in embryonated hens' eggs for virus propagation. The allantoic fluid from infected eggs was harvested after incubation of the infected eggs for 72 hrs and all the viruses were obtained from the allantoic fluid by centrifugation and were purified by sucrose gradient untracentrifugation (Skehel and Schild, 1971). The concentrations of purified viruses were determined by combined methods including gel quantification and ELISA (enzyme-linked immunosorbent assay) (Lin et al., 2012).

Biolayer Interferometry

Virus binding to defined receptor analogues was measured on an Octet RED biolayer interferometer (Pall ForteBio Corp., Menlo Park, CA, USA). Biotinylated α 2,3- and α 2,6-linked SLN were purchased from Lectinity Holding, Inc. (Moscow, Russia). These were approximately 30 kDa polymers containing 20% mol. sugar and 5% mol. biotin linked to a polyacrylamide backbone. The polymers were immobilized on streptavidin biosensors (Pall ForteBio Corp., Menlo Park, CA, USA) at concentrations ranging from 0.01 to 1.5 µg/ml. The relative sugar loading (RSL) of the biosensor was calculated from the amplitude of the response at the end of the 5-10 minute loading step. The maximum response at complete saturation was ~ 0.6 nm.

Binding of viruses (at 1nM) was measured at 25° C in a 30-50 minute association step. The buffer was 10 mM HEPES (pH 7.4), 150 mM NaCl, 2 mM CaCl₂ and 0.005% Tween-20. All solutions also contained 10 μ M oseltamivir carboxylate (Roche, Welwyn Garden City, U.K.) and 10 μ M zanamivir (GSK, Stevenage, U.K.) to prevent cleavage of the receptor analogues by the viral neuraminidase.

The (relative) amount of virus bound to the biosensor at different relative sugar loadings was calculated from the amplitude of the response at the end of the association step. The measured amplitudes were normalized by dividing by the maximum response (typically 5-6 nm) and the normalized response was plotted as a function of the relative sugar loading. The normalized virus binding response curves report the fractional saturation of the sensor surface (f) and smooth lines through the curves were generated by fitting the data to a simple variant of the Hill equation:

$$f = \frac{\left[RSL\right]^{n}}{\left[RSL_{0.5}\right]^{n} + \left[RSL\right]^{n}}$$

where RSL is the relative sugar loading, $RSL_{0.5}$ is the relative sugar loading at half saturation (f = 0.5), and n is a Hill coefficient.

Virus Preparation for Glycan Array

All viruses were inoculated in embryonated hens' eggs for virus propagation. The allantoic fluid from infected eggs was harvested after incubation of the infected eggs for 72 hrs and all the viruses were obtained from the allantoic fluid, inactivated with beta-propiolactone, and concentrated and purified by sucrose gradient ultracentrifugation. The HA titer was determined and samples were stored at -80°C.

Glycan Array Fabrication

Glycan arrays were prepared as previously described (Peng et al., 2017). Briefly, a library of asialo and sialylated glycans bearing a reducing-end amine were dissolved to 100uM concentration in printing buffer (150mM sodium phosphate + 0.005% Tween-20, pH 8.2) and aliquoted into 384-well microtiter plates. Using a MicroGrid II (Digilab) microarray printing robot, equipped with SMP3 microarray pins (Telechem), 6 replicate spots of each glycan were immobilized onto NHS-activated glass microscope slides (SlideH,



Schott). Following printing, slides were blocked to quench remaining NHS chemistry in blocking buffer (50mM ethanolamine in 50mM borate buffer, pH 9.4) and stored at -20C until used.

Virus Labeling and Binding Assays

Viruses were directly labeled with a biotin handle as previously described (Watanabe et al., 2014). Labeled viruses were diluted to 256 hemagglutinating units (HAU) in 1X PBS and applied directly to the slide surface for 1h. Following the initial incubation, arrays were washed, by dipping 3 times in 1X PBS and again 3 times in 1X PBS. Washed arrays were incubated with 2ug/mL streptavidin-Alexa-Fluor555 (LifeTechnologies) in 1X PBS, for 1h. Following detection, arrays were washed sequentially, by dipping, 3 times in 1X PBS, 3 times in 1X PBS and, finally, 3 times in deionized H2O. Washed arrays were dried by centrifugation and immediately scanned for AlexaFluor555 signal on an Innoscan 1100AL (Innopsys) confocal microarray scanner. Signal intensity from scanned arrays was collected using Mapix (Innopsys). Signal intensity was calculated for the mean signal intensity of 4 replicate spots for each printed glycan and graphed using Excel (Microsoft).

CD Measurements

Far-ultraviolet (UV) circular dichroism (CD) spectra were recorded on a Jasco J-815 spectropolarimeter fitted with a cell holder thermostatted by a CDF-426S Peltier unit. All CD measurements were made in 10mM HEPES, 150mM NaCl with and without 2 mM CaCl₂ using fused silica cuvettes with 1-mm path length (Hellma, Jena, Germany). CD intensities are presented as the CD absorption coefficient calculated on a mean residue weight basis ($\Delta \in MRW$).

Thermal unfolding curves were obtained by monitoring the ellipticity at 222 nm using 1- or 2-mm path length cuvettes and a heating rate of 1 °C/min over the temperature range 20 to 90 °C. The transition mid-point temperature was obtained by fitting to a modified Gibbs-Helmholtz equation using in-house software as described elsewhere [1].

Trp Fluorescence Assay

0.25 mg/ml HA protein in the presence or absence of 2 mM calcium was denatured using different concentrations of urea (1-10 M) at room temperature overnight except HAs with high stability that required higher temperature (30°C) to be unfolded. Fluorescence spectra were recorded on a Jasco FP-6300 spectrofluorometer. Each sample was placed into a 3 mm Cuvette and excited using 280 nm UV light and the emission spectra (from 300 to 450 nm) were obtained. The optical signals at 341nm from HAs in different concentrations of urea had maximum variation, so they were chosen to be plotted against the corresponding urea concentrations for data analysis.

Calcium Binding Assays

Macroscopic calcium binding constants (K_1 - K_3) were determined from calcium titrations performed in the presence of the chromophoric calcium chelator, 5,5'-Br₂BAPTA (5,5'-dibromo-1,2-bis(2-aminophenoxy)ethane-N,N,N',N'-tetraacetic acid), using a method described by Linse et al. [2].

Measurements were performed at 20 °C in calcium free buffer (Chelex-100 treated 10 mM HEPES, 100 mM NaCl, pH 8); under these conditions, the calcium binding constant of 5,5'-Br₂-BAPTA was determined to be 5.7×10^5 M⁻¹. The values for the individual binding constants (K1-K3) were obtained from least-squares fits directly to the experimentally observed titration curves using inhouse software.

pH Dependent Proteolysis Experiment

0.2 mg/ml HA in PBS buffer was adjusted to pH ranging from 6.84 to 4.98 using 150 mM citrate buffer, pH 3.5, and incubated at room temperature for 10 min. The pH was then adjusted to 7.5 using 1 M tris buffer, pH 8.0. The samples were treated with trypsin (10µg/ml) at 37°C for 10 min and then treated with trypsin inhibitor to stop all the reactions.

Minigenome Assay

A model vRNA, consisting of the firefly luciferase open reading frame flanked by the noncoding regions (NCRs) of segment 8 of influenza A virus, under the control of a T7 RNA polymerase promoter was used for minigenome assays (de Wit et al., 2010). The reporter plasmid (0.5 µg) was transfected into 293T cells in 6-well plates, along with 0.5 µg of each of the pHW2000 plasmids encoding PB2, PB1, PA, and NP; 1 µg of pAR3132 expressing T7 RNA polymerase (Dunn et al., 1988); and 0.02 µg of the *Renilla* luciferase expression plasmid pRL (Promega, Leiden, Netherlands) as an internal control. Forty-two hours after transfection, luminescence was measured using a Dual-Glo Luciferase Assay System (Promega) according to the instructions of the manufacturer in a TECAN Infinite F200 machine (Tecan Benelux bv, Giessen, Netherlands). Relative light units (RLU) were calculated as the ratio of firefly and *Renilla* luciferase luminescence.

Plaque Assay

The assay was performed as described (Matrosovich et al., 2006). In brief, MDCK cells (10^6 per well) were seeded in a 6 well plate to reach 90% confluency the next day. Recombinant virus supernatants were diluted and 100 µl of the selected dilution, to obtain a plaque density of ~20, was added to each well. After incubation for one hour at 37 °C and 5% CO2 cells were washed with PBS once and 2 ml of an overlay containing 2x EMEM (Lonza) and avicel (FMC BioPolymer, Newark, US) in a 1:1 ratio was added. Plates



were incubated at 37 °C and 5% CO2. After 40 h, cells were washed with PBS twice and 1 ml of 80% acetone was added. Plates were incubated at -20 °C overnight and virus infection was determined by NP antibody staining. Briefly, NP monoclonal antibody (IgG2a, clone Hb65, American Type Culture Collection, Wesel, Germany) and goat-anti-mouse Ig FITC (BD biosciences, USA) antibody were used to detect NP positive cells. Digital images were taken using ImageQuant TL software (GE Healthcare Life Sciences). The plaque sizes of some viruses were too small to analyze with the software.

Replication Kinetics in Seal Cells

Seal (*Phoca vitulina*) kidney cells (Osterhaus et al., 1985) were inoculated with 0.1 TCID₅₀/cell of virus. Cells were washed with PBS one hour after inoculation and cultured in 200 μ l of infection media, consisting of EMEM supplemented with 100 U/ml penicillin, 100 μ g/ml streptomycin, 2 mM glutamine, 1.5 mg/ml sodium bicarbonate, 10 mM HEPES, non-essential amino acids, and 20 μ g/ml TPCK treated trypsin (Lonza). Supernatant samples were harvested 6, 12, 24, 48 and 72 h later. Virus titers were determined by end-point titration in MDCK cells.

Enzyme-Linked Lectin Assay

The enzyme-linked lectin assay (ELLA) was performed as described previously to measure neuraminidase activity (Westgeest et al., 2015). In brief, fetuin (25 ug/ml) was used to coat Nunc-ImmunoTM MicroWellTM 96 well solid plates (100 μ l/well; Sanbio, Uden, The Netherlands) at 4 °C for at least 24 h. Twofold serial dilutions of reassortant viruses consisting of seven gene segments of influenza virus A/PR/8/34 and the N7 NA segment of interest were added to duplicate fetuin-coated plates. Plates were sealed and incubated at 37 °C for 16–18 h. The plates were washed and 100 μ l/well of horseradish peroxidase-conjugated peanut agglutinin lectin (PNA-HRPO, Sigma–Aldrich) was added. Plates were incubated at room temperature for 2 h. *O*-Phenylenediamine dihydrochloride (OPD, Sigma–Aldrich) substrate was freshly prepared following the manufacturer's instruction and added to the plate (100 μ l/well). The reaction was stopped after 10 min by the addition of stop solution (0.5 M H₂SO₄, 100 μ l/well). The plates were read at 490 nm (OD₄₉₀) for 0.25 s using an Infinite 200 96-well plate reader (Tecan, Giessen, The Netherlands). Next the NA activity was plotted against the antigen dilutions.

Fusion (Acid Stability) Assay

Membrane fusion was tested as previously described (Herfst et al., 2008) in a cell content mixing (CM) assay in which two 10 cm cultures dishes containing Vero-118 cells were transfected with 5 μ g of pCAGGS-HA and 5 μ g of pEGFP-N1 (as transfection control) using Xtremegene transfection reagent (Roche). One day after transfection, cell populations were harvested using trypsin-EDTA (Ethylenediaminetetraacetic acid), and plated in a 12-well plate format. The next morning, cells were exposed to PBS at different pH for 10 minutes. Cells were fixed 24 hours after the pH-pulse using 70% ice-cold acetone, washed and stained using a 20% Giemsa mixture for microscopy (Merck Millipore, Darmstadt, Germany).

QUANTIFICATION AND STATISTICAL ANALYSIS

For the ferret transmission experiments, the Fisher's Exact Test (p = 0.04) was used for the comparison of transmissibility of H10sealNL virus (transmission in 6 out of 8 ferret pairs) and H10mall (transmission in 1 out of 8 transmission pairs), Figures 1 and S1. Fluorescence intensities recorded using Mapix (Innopsys) for glycan microarray experiments in Figure 2 were quantified via measurement of mean intensity minus mean background of the four median out of six total replicate spots. Data presented is the average of these four replicates with standard error. Statistical analysis is not applied. The statistical analysis to validate all the crystal structures is included in Table S2 (Crystallographic data and refinement statistics).