Mutations in haemagglutinin that affect receptor binding and pH stability increase replication of a PR8 influenza virus with H5 HA in the upper respiratory tract of ferrets and may contribute to transmissibility

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INTRODUCTION

Highly pathogenic avian influenza (HPAI) virus H5N1 has circulated widely in a diverse range of avian species for nearly a decade. It has also caused disease in several mammals, including humans, often with lethal consequence. The current human mortality rate for infection with HPAI H5N1 is ~58% in WHO confirmed cases (WHO, 2012b). To date no sustained human to human transmission of the virus has been observed, with most human cases resulting from direct contact with sick birds (Aditama et al., 2012; Ungchusak et al., 2005; Wang et al., 2008). However, the concern remains that, if human transmissibility was acquired, a severe pandemic could result.

Vaccination is the most effective method for controlling a newly emerged influenza virus in the community. Inactivated influenza vaccines for a newly emerged subtype such as H5 to which the human population is not primed may require high antigen doses, two or more immunizations, and/or adjuvants (Couch et al., 1997; Subbarao et al., 2006). Live attenuated vaccines (LAIVs) would allow greater vaccine coverage because more doses can be generated from each infected egg. However, efforts to trial LAIVs for H5 in humans have been largely unsuccessful so far, with poor antibody responses (Karron et al., 2009). This can be explained by the cold-adapted (ca) genetic backbone of the LAIV driving virus replication to the cooler temperatures of the upper respiratory tract (URT), but the avian-adapted H5 haemagglutinin (HA) protein having low affinity for the α2,6 linked sialic acid (SA) receptors that predominate there.
Indeed, a prerequisite for an avian influenza virus to acquire transmissibility between mammals is the switch in SA linkage preference of HA from $\alpha 2,3$ linked SA to $\alpha 2,6$ linked SA, which will allow greater replication in the URT, the site of transmission. We and others have described mutations that might mediate this switch (Auewarakul et al., 2007; Ayora-Talavera et al., 2009; Chutinimitkul et al., 2010; Gambaryan et al., 2006; Gao et al., 2009; Harvey et al., 2004; Maines et al., 2011; Su et al., 2008; Watanabe et al., 2011; Yamada et al., 2006). Several groups have engineered recombinant H5N1 viruses that contain $\alpha 2,6$ SA-adapting mutations in the HA and tested transmissibility in ferrets, the best experimental animal model to indicate the potential for human influenza transmission (Chen et al., 2012a; Harvey et al., 2004; Maines et al., 2011). However, receptor binding switches alone may not mediate a complete switch of host range for H5N1 viruses. Indeed, Maines et al. (2011) showed that receptor switching mutations at the receptor binding site, Q226L and G228S, in H5 HA resulted in lower viral shedding from inoculated ferrets, and transmission remained as inefficient as for wild-type H5N1 viruses (Maines et al., 2011). However, Chen et al. (2012a) showed that insertion of three receptor binding mutations 196R/226L/228S into a clade 2.2 H5N1 virus combined with replacement of the avian virus NA (neuraminidase) gene by a human H3N2 virus NA resulted in some respiratory droplet transmission in this model. Very recently two reports of mammalian transmissible H5N1 viruses indicated a requirement for receptor binding changes in the H5 HA to increase the $\alpha 2,6$ SA affinity in combination with other changes in the HA protein (Herfst et al., 2012; Imai et al., 2012).

Wang et al. (2010) investigated whether receptor binding mutations in HA can be used to increase the nasal replication of H5 LAIV. They introduced the Q226L and G228S mutations in combination with loss of the glycosylation site at N158 and showed increased replication in ferrets. However, protection mediated by this LAIV was not cross-protective for viruses that retained the HA glycosylation site at residue 158.

One property that has been somewhat overlooked until now, which may affect the ability of avian influenza viruses to replicate in the mammalian respiratory tract, is the pH stability of the HA protein. HA is the fusogenic protein of the virus. After entry of the virus particle to the host cell, in the acidified environment of the endosome, HA undergoes an irreversible conformational change that exposes the hydrophobic fusion peptide and initiates membrane fusion leading to release of the viral genome into the cytoplasm (Skehel & Wiley, 2000). The pH at which HA undergoes this transformation varies between different strains of the virus (Beyer et al., 1986). During adaptation to growth in cell culture or to different host species, the pH at which HA fusion occurs is altered (Giannechini et al., 2006; Lin et al., 1997). Viruses bearing HA protein that fuses at a high pH release their genomes more rapidly upon cell entry and this can bestow a replicative advantage under certain conditions. However, viruses with HAs that require a lower pH to fuse display increased environmental stability and may transmit more efficiently as a result (Reed et al., 2010). The way this balance plays out for any individual virus is likely influenced by the environmental factors encountered during transmission events. For example, transmission in overcrowded poultry houses or by virus shedding into brackish water by ducks may select for HA proteins with a high pH of fusion, whereas transmission through the air into the relatively acidic environment of the human nasal cavity (England et al., 1999; Washington et al., 2000) might require a more pH-stable HA. Indeed, the ability of H5N1 viruses to replicate in the mouse URT was shown by Krenn et al. (2011) to be enhanced by mutations that stabilized the HA. We therefore speculated that mutations that lowered the pH of fusion of H5 HA might enhance its replication in the upper airways of ferrets, inducing a more robust immune response and concomitantly lead to a transmissible phenotype, at least in combination with other HA changes that affect receptor specificity and NA changes that affect virus release.

RESULTS

Human-adapted HA proteins fuse at lower pH than avian virus HAs

We assessed a panel of viruses possessing the HA and NA genes of either human transmissible viruses or avian influenza isolates (see Table 1) for the pH at which the HAs triggered membrane fusion. This was achieved by mixing 64 HA units of each virus with human red blood cells (hRBCs) at 4°C. After binding between HA and SA on the erythrocytes had taken place, the mix was exposed to gradually decreasing pH at 37°C. Fusion of membranes triggered by the HA conformational change was measured by the release of haemoglobin from the erythrocytes. The human transmissible virus HAs required a lower pH (<pH 5.3) to undergo fusion in comparison to the avian virus HAs (<pH 5.5) (Fig. 1a).

A single point mutation lowers the pH of fusion of H5 HA

We generated a panel of recombinant influenza viruses possessing the H5 HA (A/Vietnam/1194/2004) into which well characterized humanizing receptor binding site mutations were engineered, Q226L and G228S alone or coupled with the mutation T160A, which abrogates glycosylation at residue 158 (Chutinimitkul et al., 2010; Gao et al., 2009; Wang et al., 2010). We then additionally engineered another HA mutation, H24Q, which has been documented to lower the pH of fusion (Reed et al., 2009, 2010). We paired these mutated H5 HAs with either the matched A/Vietnam/1194/2004 N1 NA gene or the N1 NA gene from a prototypic isolate of the 2009 H1N1 human pandemic virus (A/England/195/2009) since this NA clearly supports efficient transmission in humans and could be acquired during a reassortment with H5N1 viruses (see...
Table 1. Genetic composition of viruses used in the study

<table>
<thead>
<tr>
<th>Virus name</th>
<th>H5 HA gene mutation (A/Vietnam/1194/2004)</th>
<th>NA gene</th>
<th>Phenotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>H5-wt*</td>
<td>Wild-type</td>
<td>A/Vietnam/1194/2004</td>
<td>Wild-type</td>
</tr>
<tr>
<td>H5-LS*</td>
<td>Q226L/G228S</td>
<td>A/Vietnam/1194/2004</td>
<td>↑≤2,6 SA binding</td>
</tr>
<tr>
<td>H5-ALS*</td>
<td>T160A/Q226L/G228S</td>
<td>A/Vietnam/1194/2004</td>
<td>↑≤2,6 SA binding</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Virus name</th>
<th>HA gene</th>
<th>NA gene</th>
<th>Phenotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>VN04 (H5 wt)*</td>
<td>A/Vietnam/1194/2004</td>
<td>A/Vietnam/1194/2004</td>
<td>Avian H5N1 (minus MBS)</td>
</tr>
<tr>
<td>HK97*</td>
<td>A/Hong Kong/156/97</td>
<td>A/Hong Kong/156/97</td>
<td>Avian H5N1 (minus MBS)</td>
</tr>
<tr>
<td>TKY05*</td>
<td>A/turkey/Turkey/01/2005</td>
<td>A/turkey/Turkey/01/2005</td>
<td>Avian H5N1 (minus MBS)</td>
</tr>
<tr>
<td>RD3*</td>
<td>A/chicken/Italy/13474/99</td>
<td>A/chicken/Italy/13474/99</td>
<td>Avian H7N1 (minus MBS)</td>
</tr>
<tr>
<td>B/Flor</td>
<td>B/Florida/04/2006</td>
<td>B/Florida/04/2006</td>
<td>Human influenza B</td>
</tr>
<tr>
<td>VIC</td>
<td>A/Victoria/3/75</td>
<td>A/Victoria/3/75</td>
<td>Human H3N2</td>
</tr>
</tbody>
</table>

Table 1). For all of the H5 recombinant viruses we generated, we removed the multi-basic cleavage site [a known major virulence factor (Hatta et al., 2001)] from the HA gene, and combined the H5 HA and N1 NA genes with the remaining six genome segments from the high growth reassortant vaccine donor strain of influenza virus, A/PR/8/34. Thus, our H5 recombinant viruses were similar in genetic makeup to the reverse genetics (RG) vaccines that have been used widely to generate and test pre-pandemic H5 inactivated vaccines (WHO, 2012a). Such viruses have been extensively tested in chickens, mice and ferrets, and demonstrated to lack the highly pathogenic phenotype [reviewed by Horimoto & Kawaoka (2009)].

We tested the effect of the engineered HA mutation (H24Q) on the pH of fusion. The human H3N2 virus (A/Victoria/3/75) did not undergo efficient fusion until exposed to a pH of <5.0 in line with the other human viruses tested in Fig. 1(a). In contrast, membrane fusion of the H5-wt virus was triggered at a pH of 5.7. The threshold pH at which fusion first occurred was not affected by the receptor specificity of the HA or by switching the avian-adapted N1 NA for the NA from the 2009 pH 1N1 virus (compare H5-wt virus with H5-ALS-pN1 in Fig. 1b). The single amino acid change H24Q lowered the H5 HA pH of fusion, such that fusion was not detected at a pH of 5.3 or above.

**pH of fusion does not affect multicycle replication in MDCK cells**

We next measured the multicycle replication of the recombinant viruses in MDCK cells. In this system all the recombinant viruses replicated robustly with released titres increasing between 24 and 48 h post-infection (p.i.). We did not observe any replicative advantage conferred by the H24Q pH stabilizing mutation in HA (Fig. 2). The combination of receptor binding changes in H5 HA with the NA gene from the pH 1N1 2009 virus (H5-ALS-pN1), did result in some enhancement of virus release at 24 h p.i. (P<0.005), likely due to an improvement in the balance between HA receptor binding and NA receptor destroying activity in MDCK cells. However, when the H24Q pH stabilizing mutation was added to this genetic constellation, the in vitro replicative advantage was tempered.

**A virus with a lower pH of fusion of HA replicates to higher titres in the ferret URT and transmits to a contact animal**

We then ascertained that experiments using a PR8 internal gene constellation with human-adapted HA and NA genes could detect efficient infection and transmission in the ferret model. Following intranasal inoculation with 10^6 p.f.u. of a recombinant virus with HA and NA gene segments of the human H3N2 virus (A/Victoria/3/75) on a PR8 internal genetic backbone, two donor animals were robustly infected. These animals shed infectious virus from the nose for the following 6 days, at titres in excess of 10^3 p.f.u. ml^-1 (Fig. S1, available in JGV Online). Moreover, the virus was transmitted to five out of six co-housed sentinel animals who themselves shed virus (Fig. S1).

We then assessed the replication of the H5 HA engineered viruses in the URT of ferrets and also tested whether the
calculated for each virus. The means of quadruplicate tests were percentage of maximum fusion achieved at each pH was the release of haemoglobin by measuring the optical density at differing pH conditions and the degree of fusion monitored by the total of 64 HA units of each virus was mixed with hRBCs in tested for their ability to cause fusion in hRBCs at varying pHs. A

Fig. 1. Influenza viruses with human- or avian-adapted HA were tested for their ability to cause fusion in hRBCs at varying pHs. A total of 64 HA units of each virus was mixed with hRBCs in differing pH conditions and the degree of fusion monitored by the release of haemoglobin by measuring the optical density at 405 nm. Each virus was tested in quadruplicate and the percentage of maximum fusion achieved at each pH was calculated for each virus. The means of quadruplicate tests were plotted with sos indicated. (a) Comparison between human-adapted influenza strains E195 - A/England/195/2009 (pH 1N1), B/Flor - B/Florida/4/2006 and SYD - A/Sydney/5/1997 (H3N2) (grey lines), and viruses bearing the surface antigens of HPAI avian viruses from which the multi-basic site had been removed from the HA on an internal genetic background of A/PR/8/34 virus as follows: VN04 - A/Vietnam/1194/04 (H5N1), TKY05 - A/turkey/Turkey/01/05 (H5N1), HK97 - A/Hong/Kong/156/97 (H5N1) and RD3 - A/chicken/Italy/13474/99 (H7N1) (black lines). (b) H5 HA mutant viruses with the multi-basic site engineered away from H5 HA and on an internal genetic background of A/PR/8/34 (H5-ALS-pN1 and H5-QALS-pN1) (black lines) were compared to H5-wt (black line) and a human virus (VIC - A/Victoria/3/75 (H3N2) (grey line). Table 1 shows the details of the genetic composition of each virus.

Mutations affected the ability of the virus to pass from one ferret to another under direct contact conditions. These experiments were performed with the knowledge and approval of local and national safety committees including the Health and Safety Executive of the UK, under containment level 3 conditions, with ferrets housed in isolator cages and all persons involved in their care or in sampling procedures donning personal protective equipment including personal respirators.

The peak titre of infectious virus shed by donor animals infected with \(10^6\) p.f.u. of the recombinant virus H5-LS, which contained the Q226L and G228S mutations, was

very low \([<100 \text{ p.f.u. (ml nasal wash)}^{-1}]\) and no onwards transmission occurred even though the sentinel animals were co-housed (Fig. 3a, Table 2). Adding the T160A mutation and coupling this HA with the NA gene from a pH 1N1 virus (H5-ALS-pN1) resulted in inoculated ferrets shedding higher titres of virus, reaching \(4 \times 10^6\) p.f.u. (ml nasal wash)\(^{-1}\) at day 1–2 post-inoculation (Fig. 3b, Table 2). However, still no transmission to naive contact animals was observed by viral detection in daily nasal wash samples and neither of the contact animals showed H5 seroconversion 21 days after exposure (Table 2).

The introduction of the single point mutation H24Q resulted in robust infection of both donor animals with titres of shed virus achieving \(2 \times 10^7\) p.f.u. (ml nasal wash)\(^{-1}\) (Table 2). Moreover, there was virological and serological evidence of transmission to one of the two co-housed sentinel animals (Fig. 3c).

**Antibodies in sera from animals infected by the mutated H5 viruses cross-neutralize lentiviral pseudotypes with wild-type H5 HA**

Sera from animals infected with H5 viruses were tested for their ability to neutralize lentiviruses pseudotyped with H5 HA from either the clade 1 A/Vietnam/1194/2004 virus, VN04 (H5-wt), or the clade 2.2.1 A/turkey/Turkey/01/2005 virus (TKY05). The lentivirus pseudotype neutralization assay is a well-accepted and safe methodology for this purpose (Alberini et al., 2009; Temperton et al., 2007). As a positive control, a ferret serum was obtained from Diane Major, National Institute for Biological Standards and Control (NIBSC), South Mims, UK, from ferrets infected with either wild-type A/Vietnam/1194/2004 virus (F1/04) or the attenuated vaccine virus RG14 which bears A/Vietnam/
H5 virus led to the generation of robust titres of antibodies by viruses with H5 HA that is more pH stable than wild-type viruses with either of the wild-type avian H5 HAs. These QALS were all able to neutralize entry of pseudotyped with humanizing receptor mutations, including LS, ALS and 3. Sera from animals infected with any of the H5 viruses neutralization activity against either H5 pseudotype (Table 3). In contrast, serum from a negative control ferret that was exposed but uninfected by pH 1N1 2009 virus (F100) had no contrast, serum from a negative control ferret that was housed individually with a naive contact ferret (sentinel). All ferrets were nasal washed daily and viral titre determined by plaque assay on MDCK cells.

1194/2004 H5 HA with a modified HA cleavage site (F5/09). These sera were effective for neutralization of the homologous and heterologous H5-pseudotyped viruses (Table 3). In contrast, serum from a negative control ferret that was exposed but uninfected by pH 1N1 2009 virus (F100) had no neutralization activity against either H5 pseudotype (Table 3). Sera from animals infected with any of the H5 viruses with humanizing receptor mutations, including LS, ALS and QALS, were all able to neutralize entry of pseudotyped viruses with either of the wild-type avian H5 HAs. These observations show that the increased replication in the nose by viruses with H5 HA that is more pH stable than wild-type H5 virus led to the generation of robust titres of antibodies that were able to efficiently neutralize wild-type H5 viruses of a different clade.

DISCUSSION

Recently two laboratories showed that ferret-passaged highly pathogenic H5N1 influenza virus acquired aerosol transmissibility in ferrets following mutations in the HA gene that both enhanced the HA affinity for α2,6 linked SA and affected the stability of the HA molecule (Herfst et al., 2012; Imai et al., 2012). The viruses utilized for those two studies retained genetic characteristics associated with high virulence, for example, the multi-basic cleavage site of H5 HA and, in the case of the work from Herfst and colleagues, an internal genetic constellation derived from the H5N1 virus that may be associated with induction of a cytokine storm (Mok et al., 2009). The work we describe here shares some similarities in approach with those two studies in that we have engineered humanizing mutations into influenza viruses of the H5 subtype. However, we chose to perform this work in a virus genetic background where the HA pathogenicity determinant (multi-basic cleavage site) was removed and the internal genes were replaced by those of the PR8 high growth reassortant vaccine donor strain, which has a long history of safe use. This undoubtedly led to a decrease in the levels of virus titres accumulated in the nose of infected ferrets compared with the wild-type genetic background. Indeed, a comparison of nasal virus titres shed from ferrets infected with the VIC/PR8 virus in Fig. S1 with titres on equivalent days shed from ferrets infected with a wild-type Victoria virus (Roberts et al., 2011) indicated that the internal genes of PR8 reduced virus shedding by 2–3 logs. Thus, we do not suggest that we would measure respiratory droplet transmission using this genetic backbone, which is a stringent measure of mammalian adaptation. Nonetheless, our data suggest similar conclusions to those from the work with HPAI virus, that increased replication and consequent onwards transmission of H5 influenza virus may be increased in ferrets by combinations of mutations that affect at least three biological traits: alteration of receptor binding preference, stabilization of HA and a replication-competent polymerase (Imai et al., 2012). In addition, our results suggest that a NA gene adapted to human transmission may enhance URT shedding and onwards transmissibility (Fig. 2). We did not formally establish the effect of the NA gene in this study, by comparing infection and transmissibility of, for example, an H5-ALS mutant possessing the Vietnam N1 NA with that of a virus containing the same HA coupled with pH 1N1 NA. However, several other studies have suggested a role for NA in transmission (Chen et al., 2012a; Lakdawala & Subbarao, 2012; Lakdawala et al., 2011) and our own studies had reinforced this (D. Blumenkrantz & W. S. Barclay, unpublished data). Therefore, we chose to use the pH 1N1 NA in the present studies in order to allow us to focus on the effects of changes in the H5 HA.

Of these proposed adaptations, the requirement for pH stabilization is the most novel. The pH at which HA fusion
occurs is a compromise between the need for virus to survive in the environment and the enhancement to infectivity that can be conferred by fusion and uncoating in early rather than in late endosomes. This balance will be altered when the virus crosses from one species to another and finds a new environmental niche across which transmission must occur. The fusion pH of HA has already been shown to be important in the transmission of H5N1 viruses in ducks. In this natural avian host, mutations that stabilized HA, decreasing the pH of fusion below pH 5.6, resulted in a reduction in bird to bird transmission (Reed et al., 2010). It was also recently shown that the pathogenicity of HPAI H5N1 in chickens is affected by the pH of HA fusion, with an increased pH of fusion leading to a more pathogenic phenotype in vivo (DuBois et al., 2011). In our study, we did not find an effect of NA on the pH of HA fusion, rather the HA mutation in the stalk region of HA1 subunit had the dominant effect.

The mutation we used to stabilize H5 HA, H24Q, was introduced into the H5 HA of A/Vietnam/1194/2004, a clade 1 virus for which a licensed pre-pandemic vaccine, (Prepandrix; GlaxoSmithKline Biologicals) has been generated and for which a crystal structure is available (Stevens et al., 2006; Yamada et al., 2006). This residue is located in the fusion peptide pocket of the H5 HA, where it is one of three conserved histidines that at low pH become positively

<table>
<thead>
<tr>
<th>Virus name</th>
<th>Infected</th>
<th>Peak viral titre shed (p.f.u. ml⁻¹)</th>
<th>Day of viral peak</th>
<th>Viral shedding</th>
<th>Seroconversion</th>
</tr>
</thead>
<tbody>
<tr>
<td>H5-LS*</td>
<td>2/2</td>
<td>35</td>
<td>3</td>
<td>0/2</td>
<td>0/2</td>
</tr>
<tr>
<td>H5-ALS-pN1*</td>
<td>2/2</td>
<td>15</td>
<td>1</td>
<td>0/2</td>
<td>0/2</td>
</tr>
<tr>
<td>H5-QALS-pN1*</td>
<td>2/2</td>
<td>3 × 10⁴</td>
<td>3</td>
<td>0/2</td>
<td>0/2</td>
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<td></td>
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<td>4 × 10⁴</td>
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<tr>
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<td></td>
<td>1.75 × 10⁵</td>
<td>1</td>
<td></td>
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</table>

*Viruses contained the six internal gene segments from A/PR/8/34.
charged and cause the fusion peptide to exit the pocket in an irreversible conformational change (Stevens et al., 2004). Reed et al. (2009) already demonstrated that the H24Q mutation reduced the pH of fusion of a closely related H5N1 virus (A/chicken/Vietnam/CS8/2004). However, there are several other genetic changes by which the H5 HA might acquire increased pH stability. Interestingly, during the independent studies involving ferret passage leading to transmission of recombinant H5N1 influenza viruses, different mutations in the HA stalk region were selected that resulted in increased pH stability (Herfst et al., 2012; Imai et al., 2012). Mutations that H5N1 viruses are accumulating whilst circulating in the wild bird population are currently monitored for changes to the receptor binding site of the HA protein (Watanabe et al., 2011). It now seems important to also monitor for changes that affect pH stability so that viruses with increased pandemic potential that arise in nature can be detected early.

Ideally, during this pre-pandemic grace period we will improve our ability to generate effective vaccines against novel influenza subtypes. LAIVs for seasonal human influenza are used widely in some countries, especially in children, an important target group who are major spreaders of virus through the community (Carrat et al., 2006; Cauchemez et al., 2011; Loeb et al., 2010). However, pre-pandemic avian influenza LAIVs currently induce only a poor immune response due to the conflict between their restriction to the URT by cold-adapted internal genes and the lack of α2,6 SA tropism of their HA. Moves to increase the ‘take rate’ of LAIVs by altering receptor binding or lowering the pH of fusion have both been explored separately (Krenn et al., 2011; Wang et al., 2010). Here, we combined these two approaches and showed an additive effect. It may be useful to use one or other strategy separately to generate an H5 HA LAIV with higher nasal replication but which is still unable to transmit onwards to non-immunized individuals. Most important though, is the assessment of whether by engineering mutations into the H5 HA that increase nasal replication there is any unforeseen effect on antigenicity that might compromise the efficiency of the induced immune response to protect vaccinees. Wang et al. (2010) made H5N1 LAIV with humanizing mutations in the HA receptor binding site, including deletion of the glycosylation site at residue 158, and found that the sera generated in ferrets infected with these vaccine candidates were inefficient at recognizing the wild-type H5 HA (Wang et al., 2010). This may be because glycosylation of T158 interferes with antibody access to an important epitope (Bender et al., 1999). Similarly Chen et al. (2012b), striving to increase the vaccine immunogenicity of LAIV against H2 and H6 avian subtypes of influenza by engineering humanizing mutations in the receptor binding site, also found that although homologous titres were increased, the sera induced by immunization with the viruses with mutated HA did not cross-protect efficiently against challenge with virus that retained wild-type avian HA. In our study we demonstrated using ferret sera from animals infected with humanized H5 HA viruses that both receptor switching mutations and pH stability alterations led to a robust immune response, and the resulting sera could neutralize lentivirus pseudotyped with homologous wild-type H5 HA (Table 3). Moreover, in animals infected with the ‘double’ QALS mutant, a considerable titre of cross-clade neutralizing antibody titre was induced without compromising the titre against the homologous wild-type H5 HA. This suggests that even when wild-type HA retains glycosylation that may mask some antigenic epitopes, the cost to neutralizing ability can be compensated for by inducing a stronger immune response.

Since the exact nature of mutations that will ultimately arise in nature and lead to transmissible H5 virus cannot be exactly predicted, it will be important to continue research aimed to understand how mutations in receptor binding and pH stability affect the immune response generated by recombinant virus vaccines. Overall, this work has important implications for the rational design of LAIVs or more traditional pre-pandemic H5N1 vaccines, as well as for understanding the barriers that limit avian influenza virus transmissibility in mammals.

**METHODS**

**Reagents, cells and viruses.** 293T and MDCK cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM; Gibco, Invitrogen) supplemented with 10 % FBS (Biosera) and 1% penicillin/streptomycin (Gibco, Invitrogen) in 5% CO2. Wild-type human viruses, A/Sydney/2007/97 (H3N2), A/Victoria/3/75 (H3N2), A/England/195/09 pH 1N1 and B/Florida/04/06, and recombinant viruses as below were grown in MDCK cells.

**Generation of H5 HA mutant viruses.** Recombinant viruses were generated by 12 plasmid RG in 293T and MDCK cells as described elsewhere (Ellman & Barclay, 2004; Neumann & Kawaoka, 1999). Recombinant viruses possessed the six internal genes (PB1, PB2, PA, NP, M and NS) from the high growth reassortant vaccine donor strain A/PR/8/34 and the wild-type or mutated surface proteins HA and NA from the following strains: A/Vietnam/1194/04 (H5N1), A/turkey/Turkey/01/05 (H5N1), A/Hong Kong/156/97 (H5N1) or A/chicken/Italy/13474/99 (H7N1). All H5N1 and H7N1 viruses were generated from plasmids where the multi-basic site was replaced with a mono-basic cleavage site known to ablate the high pathogenicity of these HAs. Site-directed mutagenesis on the H5 HA gene from A/Vietnam/1194/04 generated receptor binding mutations (T160A, Q226L and G228S) and the pH stabilizing mutation (H24Q) in various combinations (see Table 1 for virus genetic details).

**pH of fusion assays.** Assays of the pH for HA fusion were carried out as described by Krenn et al. (2011) with the following modifications: 64 HA units per 50 μl each virus were mixed with 150 μl 1% (w/v) suspension of hRBCs in PBS and incubated at 4 °C for 1 h to allow HA attachment to SA. The virus and hRBC mix was then pelleted, washed in PBS and resuspended in MES buffer of varying pH values (from pH 4.6 to 6.0) followed by incubation at 37 °C for 60 min to allow fusion. Fusion results in release of haemoglobin from the hRBCs, which can be quantified by measuring the optical density at 405 nm. A mock control, without any virus, was performed for each pH value to control for effects of lowered pH on the stability of the hRBCs. The mock value for optical density at
405 nm for each pH value was subtracted from each virus reading, and then the percentage haemolysis was expressed as a percentage of the maximal reading for each virus. At the lowest pH of 4.6, this background level of haemolysis was not more than 20% of the raw optical density value obtained with any virus. Every virus at each pH condition was performed in quadruplicate and plotted as a mean ± SD as error bars.

**MDCK growth curves.** Confluent MDCK cells in triplicate were infected with an m.o.i. of 0.001 of each virus for 1 h at 37 °C before the inocula were removed and replaced with serum free DMEM (Gibco, Invitrogen) plus 1.4 μg TPCK trypsin ml⁻¹ ( Worthington). Virus used for inoculation was back-titrated on the day (time 0) by plaque assay on MDCK cells to confirm that inputs contained equal m.o.i. Cells were incubated at 32 °C and at 24, 48 and 72 h p.i. an aliquot of the medium was taken from each well and titrated by plaque assay on MDCK cells. Mean p.f.u. ml⁻¹ was plotted with SD as error bars. An unpaired *t*-test was used to assess the statistical significance between viruses.

**Ferret studies.** All experimental conditions were approved by local and national safety committees including the Health and Safety Executive in the UK. The experimental set up was approved also by the Home Office in the UK, who licence experimental use of animals. Female ferrets (14–16 weeks old) weighing 700–900 g were used. After acclimatization, sera were obtained and tested by micro-neutralization assay for antibodies against A/England/195/09, a prototypic 2009 H1N1 influenza virus, which was the predominant virus circulating in the community at the time of these experiments. All ferrets were negative for influenza antibodies at the start of the experiments. All ferret studies were performed in a containment level 3 laboratory with the ferrets housed in HEPA (high efficiency particulate air)-filtered isolators then nasal washed and weighed over a downdraft table. Ferrets were moved between the isolator and downdraft table in a HEPA-filtered transport box so the ferrets were under primary containment at all times except whilst on the downdraft table. All personnel involved in the care and experimental sampling of the ferrets wore personal protective equipment, including personal 3M filtered respirators. Ferret body weight was measured daily. Sentinel animals were handled before inoculated animals, and work surfaces decontaminated and handlers' gloves exchanged between animals. For inoculation, ferrets were lightly anaesthetized with ketamine (22 mg kg⁻¹) and xylazine (0.9 mg kg⁻¹), and inoculated intranasally with 10⁵ p.f.u. virus diluted in PBS (0.1 ml per nostril) on the downdraft table. Each of two inoculated donor ferrets was co-housed after 24 h with one naive sentinel animal. All animals were nasal washed daily, while conscious, by instilling 2 ml PBS into the nostrils, and the expectorate was collected in modified 250 ml centrifuge tubes. BSA (0.3 %) was added to the nasal wash expectorate before infectious virus was titrated by plaque assay on MDCK cells. The limit of virus detection in the plaque assays was 10 p.f.u. ml⁻¹. Ferrets were housed for 21 days from inoculation when they were humanely euthanized and terminal blood collected for analysis by microneutralization assay.

**Pseudotype construction and neutralization assays.** Lentiviral pseudotypes with HA glycoproteins derived from the HPAI viruses H5N1 clade 1 A/Vietnam/1194/04 and clade 2 A/turkey/Turkey/01/05 were constructed as described previously (Tempert et al., 2007). Pseudotype viruses were produced by co-transfection of 293T cells with the respective HA plasmids, the human immunodeficiency virus gag-pol plasmid p8.91 and the reporter plasmid pCSFLW (expressing firefly luciferase) using the FuGENE-6 transfection reagent (Roche). For the neutralization assay, serum samples (5 μl) were heat inactivated at 56 °C for 30 min, twofold serially diluted in culture medium and mixed with pseudotype virus [1 × 10⁶ relative light units (RLU) luciferase input] at a 1:1 (v/v) ratio. After incubation at 37 °C for 1 h, 1 × 10⁴ 293T cells were added to each well of a white 96-well flat-bottomed tissue culture plate. Firefly RLU were evaluated 48 h later by luminometry using the Bright-Glo assay system (Promega). IC₅₀ end-point titres were calculated using GraphPad Prism software from RLU values obtained from duplicate serum assays and from pseudotype virus only/cell only controls.

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