The bat influenza H17N10 can be neutralized by broadly-1

neutralizing monoclonal antibodies and its neuraminidase can 2

- facilitate viral egress. 3
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- 17 Key words: Bat Influenza virus, broadly neutralizing antibody, Hemagglutinin, Neuraminidase

Abstract 18

19 The diversity of subtypes within the Influenza A virus genus has recently expanded with the 20 identification of H17N10 and H18N11 from bats. In order to further study the tropism and zoonotic 21 potential of these viruses, we have successfully produced lentiviral pseudotypes bearing both 22 haemagglutinin H17 and neuraminidase N10. These pseudotypes were shown to be efficiently 23 neutralized by the broadly-neutralizing monoclonal antibodies CR9114 and FIG. Our studies also 24 confirm previous reports that H17 does not use sialic acid as its cellular receptor, as pseudotypes 25 bearing the H17 envelope glycoprotein are released into the cell supernatant in the absence of NA. However, we demonstrate that N10 facilitates heterosubtypic (H5 and H7) influenza HA-bearing 26 27 pseudotype release in the absence of another source of NA, significantly increasing luciferase 28 pseudotype production titres. Despite this, N10 shows no activity in the enzyme-linked lectin assay 29 used for traditional sialidases. These findings suggest that this protein plays an important role in viral 30 egress, but is perhaps involved in further accessory roles in the bat influenza lifecycle that are yet to be discovered. Thus we show the lentiviral pseudotype system is a useful research tool, and 31 32 amenable for investigation of bat influenza tropism, restriction and sero-epidemiology, without the 33 constraints or safety issues with producing a replication-competent virus, to which the human 34 population is naïve.

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35 Significance statement

36 Influenza virus is responsible for mortality and morbidity across the globe; human populations are 37 constantly at risk of newly emerging strains from the aquatic bird reservoir which harbors most of 38 the subtypes of influenza A (H1-H16). Recently identified subtypes (H17N10, H18N11) from bats 39 have broadened the reservoir from which potential pandemic strains of influenza can emerge. To 40 evaluate the potential for these novel subtypes to cross over into human populations, their ability to 41 establish an infection, in addition to the extent of cross-reactive immunity established by human 42 seasonal strains needs to be investigated. This study highlights a novel platform for the study of the 43 bat H17 and N10 envelope glycoproteins, using a lentiviral pseudotype system. Following the 44 generation of this pseudotype it was employed in cell entry and microneutralization assays. These 45 showed that two well-characterised monoclonal antibodies (mAb) which target avian and human 46 influenza subtypes will also neutralize H17. Furthermore the data presented in this study show a 47 novel aspect of the N10 glycoprotein in its ability to facilitate the budding of pseudotypes bearing 48 different influenza HAs.

49 Introduction

50 Influenza A virus is the principal causative agent of influenza, which is a substantial burden to global 51 economies and represents a significant public health risk worldwide (Ma et al. 2009; Reperant et al. 52 2012). The first stages of the viral life cycle are mediated by the influenza glycoprotein 53 hemagglutinin (HA), which is located in the lipid outer membrane of the virus. This protein mediates 54 both the attachment of the influenza virus to the target cell and the fusion process which allows the 55 virus to infect its host cell. Through its activity, HA is a key determinant of virus tropism (Dos Reis et 56 al. 2011; Sahini et al. 2010). Furthermore, HA is a highly variable protein and its features are often 57 used as methods for distinguishing different viral strains (Ebrahimi et al. 2014; Wilson and Cox 58 1990).

59 While the association between influenza viruses and wild birds has long been established (Gelfond 60 et al. 2009; Webster et al. 1992), the discovery of novel influenza-like viruses in New World bats 61 (H17N10, H18N11) represents a possible challenge to the notion that avian species are its sole 62 reservoir (Freidl et al. 2015; Sun et al. 2013; Tong et al. 2012, 2013). Recently, Egyptian *Rousettus* 63 *aegyptiacus* bats have been reported to harbor a third bat influenza subtype (Kandeil et al. 2018). It 64 is highly likely that further subtypes will be discovered in future years, warranting further research 65 on existing strains of bat influenza.

The H17N10 and H18N11 subtypes of influenza A were discovered in bat species in Peru and Guatemala respectively. They are significantly diverged from other known influenza strains, in respect to all eight gene segments (Tong et al. 2012, 2013). The HA of these novel strains contain a number of apparently unique structural features and exhibit receptor-binding activities that differ from other influenza viruses (Sun et al. 2013). Notably, it has been shown that H17 and H18 hemagglutinins do not bind to sialic acid receptors, and the true receptors are currently still unknown (Tong et al. 2013).

The neuraminidase (NA) of these viruses is also divergent from other known NAs, but the overall structure is preserved, despite differences in the active site (Juozapaitis et al. 2014; Ma et al. 2015).
Lack of sialidase activity has been reported previously for these NAs (García-Sastre 2012; Li et al. 2012; Tong et al. 2013; Zhu et al. 2012), suggesting they utilize a different substrate altogether. The combination of atypical HA binding profiles alongside a lack of NA activity suggests that this virus

functions differently to previously discovered influenza A subtypes, despite relative phylogeneticrelatedness.

Although the receptor remains unknown, H17 and H18 sequences have been extensively analyzed and compared to those of other HA gene sequences, to investigate the potential for these bat viruses becoming zoonotic (Freidl et al. 2015; Mänz et al. 2013).

At present, the degree to which the H17N10 and H18N11 isolates are capable of infecting non-bat hosts is unknown, as attempts to isolate wildtype virus have not been successful. Sequence analysis indicates that there is significant potential for a spillover occurrence, but further research is required to assess the true potential of these viruses as pandemic threats (Freidl et al. 2015; Juozapaitis et al. 2014).

To study the zoonotic potential of H17N10 and H18N11 viruses, attempts to integrate bat HAs into influenza A reverse genetics systems were made but none yielded infectious virus (Juozapaitis et al. 2014; Zhou et al. 2014). More recently, progress has been achieved by Moreira et al. 2016 to this end, using a pseudotyped VSV platform.

92 Influenza pseudotyped viruses (PVs) are useful tools to study both viral entry mechanisms and the 93 antibody response directed against the influenza HA and NA. Furthermore, when use of PVs is 94 coupled with detailed sequence analysis and phylogenetic inference, they offer the potential to 95 establish safe and effective assays to inform epidemiological and public health models of viral spread 96 and risk. This is especially the case with later generation, single cycle lentiviral vectors, allowing 97 experimentation on functional glycoproteins with the use of a reporter incorporated in the lentivirus 98 genome.

In this study, we report generation of the first H17 and H17N10 pseudotyped lentiviruses, their use
 in virus neutralization assays using broadly neutralizing monoclonal antibodies (bnmAbs) and
 investigation into the debated substrate specificity of the putative N10 neuraminidase.

102 **Results**

103 Generation of H17- and N10- pseudotype viruses

104 Generation of H17- (A/little vellow shouldered bat/060/2010) and H5- (A/Vietnam/1194/2004) 105 lentiviral PVs was achieved after transfection of HEK293T/17 cells with lentiviral pseudotype 106 production plasmids (Figure 1), followed by transduction of target human U87 MG (glioblastoma) cells with PV supernatant. Generation of H17- PVs was only possible in the presence of HAT or 107 108 TMPRSS2 proteases in HEK293T/17 cells (Figure 2), as has been seen with other subtypes of 109 influenza (Bertram et al. 2010; Böttcher et al. 2006). Subsequent experiments were carried out using 110 canine MDCK II and RIE1495 target cells which were previously reported to allow production of VSV-111 H17 (Moreira et al. 2016). In initial experiments, attempts were made to generate functional H17 112 and H17N10 PV utilizing different protease-expressing plasmids to promote HA maturation. PV 113 luciferase-based titres increased significantly when HAT-, TMPRSS2- and TMPRSS4- expression 114 plasmids were used in the transfection mix when compared to controls, indicating activation of the 115 bat HA and fusion competence. Other proteases tested (furin, KLK5, and TMPRSS3) did not yield 116 significant titre increases. MDCK II and RIE1495 cells were notably transduced, whereas HEK293T/17 117 cells were not. In the absence of co-transfected, protease-encoding plasmids, no significant PV titre 118 was measurable (Figure 3).

119 H17 entry of target cells is inhibited by broadly neutralizing monoclonal antibodies

120 Treatment of RIE1495 cells with the acidifying agent ammonium chloride resulted in significantly 121 lower levels of luciferase activity, demonstrating that H17 requires a low pH for membrane fusion 122 similar to conventional influenza viruses, such as H5 (Figure 4). Several studies have demonstrated 123 that the HA trimers of conventional influenza viruses must simultaneously coordinate their 124 conformational changes to complete membrane fusion (Otterstrom et al. 2014). HA-binding broadly 125 neutralising antibodies (bnmAbs) typically recognize loop regions surrounding the receptor binding 126 site or conserved regions of the stem and their induced-inhibition is serotype-specific (Dreyfus et al. 127 2013; Ekiert et al. 2009; Sui et al. 2009). In order to test whether cross reactive antibody responses 128 would affect the H17 glycoprotein, several characterised bnmAbs were employed. Neutralization 129 potency was measured as IC_{50} (concentration or serum dilution required to neutralise 50% of input virus). Neutralization tests were conducted using both the permissible target cell lines identified 130 131 above (MDCK II and RIE1495). The first bnmAb, CR9114, binds to a conserved epitope in the HA stalk 132 of group 1 and 2 influenza A viruses, and has also been shown to protect against lethal influenza 133 challenge in a mouse model against both lineages of influenza B (Dreyfus et al. 2012). The second, 134 CR6261, has been shown to neutralize H1 and H5 subtypes (Friesen et al. 2010). The third, FI6-nt, 135 can neutralize H1 to H16 subtypes of influenza A. A fourth, FI6-chol is a derivative of F16-nt, 136 conjugated to cholesterol (Corti et al. 2011; Lacek et al. 2014). H17 is effectively neutralized by 137 CR9114, at a concentration of 0.05 µg/ml. FI6-nt and FI6-chol also neutralize H17N10- PV, however, 138 concentrations required exceed those of CR9114. In contrast, CR6261 was not able to neutralize 139 H17 using the H17 PV assay (data not shown). Neutralization data is shown in Figure 5.

140 N10 facilitates production of conventional H5 and H7 bearing PV.

141 Pretreatment of target cells with sialidase, which removes cell surface sialic acids, did not affect the 142 entry of H17 and H17N10 PVs, which supports previous studies that H17 does not bind sialic acids 143 for infection (Maruyama et al. 2016), see Figure 6. N10-bearing PVs were titrated alongside H17 and 144 H17N10 PV, in order to measure their ability to transduce cells. Results showed that N10 had no 145 effect on H17 mediated viral entry and did not improve transduction when co-expressed with H17 146 (data not shown). However, addition of N10 in the generation of H5 or H7 PVs (H5N10, H7N10) 147 produced high titre PV in the absence of any other neuraminidase source (Figure 7), indicating that 148 N10 is facilitating release of viral particles bearing sialic acid-binding glycoproteins. H5N10 luciferase 149 titres were one log lower than parental H5 PV produced with exogenous neuraminidase, but two 150 logs higher than N10, Δ Envelope (Δ Env) or cell only controls (Figure 7). Despite this, the same 151 H5N10 PV preparation did not show any detectable NA activity when titrated using the enzyme 152 linked lectin assay (ELLA), confirming previous reports that this protein is not a sialidase (García-153 Sastre 2012; Li et al. 2012; Tong et al. 2013; Zhu et al. 2012). Similarly, multiple different 154 preparations of H17N10 or of N10 bearing PV produced negative results in ELLA (data not shown).

155 Discussion

We have successfully produced H17 and H17N10 bearing lentiviral pseudotypes, and shown that N10 is not required for H17 pseudotype budding, but that it can mediate release of a heterologous, sialic acid-binding HA bearing PVs. These bat influenza PVs were neutralized by cross-reactive bnmAbs, suggesting that the stalk region of the H17 glycoprotein retains conserved epitopes present in group 1 and 2 influenza HAs (Sun et al. 2013).

Of the target cell lines tested, H17 and H17N10 lentiviral PV were able to transduce U87 MG, MDCK II and RIE1495 cells to varying degrees when activated by proteolytic cleavage. Reports differ on the permissibility of the MDCK I type cells (Hoffmann et al. 2016; Moreira et al. 2016). Due to the nature of the pseudotype based microneutralisation (pMN) assay used in this study, the addition of cells in suspension into PV containing supernatant may allow infection/transduction to occur before cell 166 adhesion and thus by a different route than in vivo. This may explain our results in the context of 167 Moriera and colleagues' findings that the bat influenza H18 VSV pseudotypes enter at the 168 basolateral membrane and inefficiently infected confluent cell monolayers (Moreira et al. 2016). 169 MDCK I and II differ in passage number (low and high, respectively), differences between them 170 include the mucin-type transmembrane protein podoplanin, which is expressed only in MDCK I, and 171 the Forssman glycosphingolipid, which is expressed only in MDCK II cells (Hansson et al. 1986; 172 Zimmer et al. 1997). The susceptibility of MDCK cells to bat influenza viruses is unclear, and is partly 173 compounded by the availability of various lineages with different characteristics (Dukes et al. 2011). 174 The widespread use of MDCK cells in influenza research is at odds with a previous study that showed 175 H17 failed to bind to MDCK cells (Sun et al. 2013). While the reasons for this are currently unclear, it 176 has been hypothesized that the level of HA binding is below the threshold of detection for the assays 177 used in this earlier study (Maruyama et al. 2016). Alternatively, only a subset of MDCK lineages may be susceptible to these viruses (Table 2). RIE1495 cells are morphologically similar to MDCK II cells 178 179 and express the Forssman antigen, a possible factor in their susceptibility to H17 bearing PVs 180 (Moreira et al. 2016). The cell tropism data reported in the Moreira study raise some interesting 181 ideas concerning the H17 receptor, as only three out of eight bat cell lines were transduced by the 182 H17-VSV pseudotypes, originating from two *Miniopterus* and one *Pteropus* species. This suggests 183 that these viruses may be restricted to a particular set of closely related species belonging to the 184 Miniopteridae and Pteropidae families, which are closely related to the Phyllostomidae family, from 185 which the original H17 and H18 samples were isolated (Agnarsson et al. 2011; Tong et al. 2012). 186 However, this raises further questions, as the cell lines derived from *Pteropus dasymallus* 187 yayeyamae and Rousettus leschenaultiii were not permissive, indicating a complex pattern of 188 susceptibility of species to these new influenza viruses. Nevertheless, as these cell lines have only 189 recently been isolated and immortalised (Maeda et al. 2008; Maruyama et al. 2014), detailed 190 characterisation of their surface proteins and expressed proteases are not yet available, requiring 191 additional research before conclusions can be drawn. Further investigation is required to determine 192 the mechanisms involved for the transduction of canine cells but not HEK293T/17 cells, particularly 193 relating to the expression of putative receptors on permissive cell lines.

194 Our results, contrary to previous studies, indicate that N10 is performing a similar function to other 195 NAs in enabling release of new influenza virus particles, which indicate its ability to form VLPs rather 196 than releasing H17 from its cellular receptor (Yondola et al. 2011). PVs bearing H5/H7 and N10 197 envelope glycoproteins successfully budded from producer cells into the surrounding medium in the 198 absence of a sialic acid cleaving neuraminidase, resulting in significantly increased titres when 199 compared to the same glycoproteins generated without the addition of N10 (Figure 7). This increase 200 in budding PV may be due to the action of N10 on an unknown substrate, or perhaps action of the 201 protein itself in virus morphogenesis and budding (Barman et al. 2004; Yondola et al. 2011). 202 Sialidase activity was however not detected using the enzyme-linked lectin assay, suggesting either a 203 lack of sensitivity of this assay or another mechanism for the removal of surface sialic acids by N10 204 (Juozapaitis et al. 2014; Sun et al. 2013; Wu et al. 2014). Similarly, PVs bearing solely the N10 205 glycoprotein did not show sialidase activity via ELLA, or successfully transduce cells to give a 206 significant luciferase reading. Further investigation is required on the role of the bat influenza 207 neuraminidase in its lifecycle, the combination of the findings described in this article, with the fact 208 that the N10 enzymatic structure remains conserved and NA-like, suggests that it is involved in 209 accessory functions other than simply aiding in viral egress. Our results highlight the distinct 210 difference between bat and traditional influenza A viruses where a delicate balance is in place 211 between HA and NA. It may be the case that such a balance exists between bat HA and NAs which 212 will be made clear once the substrate(s) of the bat NA is discovered.

213 In a previous study, it was demonstrated that the TMPRSS2 protease was capable of inducing HA 214 maturation of H17 through cleavage from HA0 to HA1 and HA2 (Hoffmann et al. 2016). In our study, 215 we demonstrated that this maturation can also be facilitated by TMPRSS4 and HAT. This is of 216 particular interest as the expression of specific proteases is a known limiting factor in viral tropism 217 for a number of different viruses (Böttcher-Friebertshäuser et al. 2010; Ferrara et al. 2012; Millet 218 and Whittaker 2015). This, coupled with the observed susceptibility of a canine-derived cell line to 219 the H17 PV, will need to be factored into future analysis of the potential for zoonotic spillover from 220 bat origin influenza viruses.

221 The fact that bnmAbs were able to neutralize the virus particles via the H17 hemagglutinin, and their 222 requirement for proteolytic activation, reinforces that we have only scratched the surface in terms 223 of their characterization. Further exploration is required to establish whether these viruses are 224 endogenous or capable of forming infectious particles in vivo. Research has revealed that all eight 225 segments of the genome encode functional proteins (Juozapaitis et al. 2014; Moreira et al. 2016; Wu 226 et al. 2014; Zhu et al. 2012), despite the inability to isolate wildtype virus to date. Nevertheless, the 227 development of tools, such as those described here, which can be used in H17 receptor 228 identification studies may ultimately aid discovery of wildtype virus samples from the bat reservoir.

229 Methods

230 Plasmids:

The H17 HA and N10 NA genes from A/little yellow shouldered bat/Guatamala/060/2010 were 231 232 synthesised commercially by Genscript (Genscript, USA) and subcloned into vector pl.18 (Cox et al. 233 2002). Lentiviral packaging plasmids p8.91 (Zufferey et al. 1997) and pCSFLW (Demaison et al. 2002) 234 containing a firefly luciferase reporter were used to produce PV. Protease encoding plasmids 235 phCMV-Tag3 (TMPRSS4-myc) and pcDNA3.1-hTMPRSS3 were kindly provided by Prof. Stefan 236 Pöhlmann, Infection Biology Unit, German Primate Center, Germany. pCAGGS-TMPRSS2 and 237 pCAGGS-HAT were kindly provided by Eva Böttcher-Friebertshäuser, Philipps University of Marburg, 238 Germany. The plasmid bearing the Vesicular stomatitis virus envelope protein (VSV-G), pMD.G was 239 obtained from Dr Yasu Takeuchi, University College London, United Kingdom. All the plasmids that 240 were used in this study are listed in Table 3.

241 Cell lines:

Multiple cell lines were used for titration of PV containing supernatants. HEK293T/17 cells were kindly provided by Dr Edward Wright (University of Westminster, UK). MDCK II and RIE1495 were kindly provided by Dr. Gert Zimmer (Institute of Virology and Immunology, Switzerland). U-87 cells were provided by Dr Simon Scott (University of Kent, UK). Madin-Darby Canine Kidney (MDCK) cells were kindly provided by Prof. Sarah Gilbert (Jenner Institute, University of Oxford, UK). All cells were cultured in Dulbecco's modified eagle medium supplemented with 10% fetal bovine serum.

248 Antibodies:

249 mAbs CR9114 and CR6261 were produced by Crucell (Janssen Vaccines AG, Bern, Switzerland). FI6-nt

- and cholesterol conjugated FI6 (FI6-Chol) were produced by Alfredo Pesci and Krzysztof Lacek from
- 251 sequence information derived from Corti et al. 2011.

252 **Production and quantification of H17N10 and H17 bearing lentiviral pseudotypes:**

PV were produced as described previously (Ferrara et al. 2012; Temperton et al. 2007) and as shown in Figure 1. Briefly, transfection of HEK293T/17 cells was performed using a variety of combinations 255 of plasmids pl.18-H17, pl.18-N10, p8.91 and pCSFLW using polyethylenimine transfection reagent 256 (Sigma Aldrich, UK). Protease-encoding expression plasmids were also included (Table 3). Medium 257 was replaced 12h post-transfection. Supernatants were harvested 48h post-transfection and passed 258 through a 0.45µm filter (Millipore, UK). PV-containing supernatants were titrated using the firefly 259 luciferase Bright-Glo[™] system (Promega, UK). Serial dilution of 100µl of PV-containing supernatant was performed across a white flat bottom 96-well Nunclon© plate (Thermo Fisher Scientific, UK). 260 Subsequently, approximately 1×10^4 cells per well were added per well in 50µl of medium, plates 261 were incubated in a humidified incubator at 37°C 5% CO₂ for 48h, after which 50µl of Bright-Glo™ 262 263 substrate was added and luciferase reading recorded in relative luminescence units (RLU) after a 5 264 minute incubation period. Further sets of N10-bearing PV were produced by transfection of 265 HEK293T/17 cells with 500ng p8.91, 750ng pCSFLW and various amounts of pl.18-N10 plasmid. 266 Medium was replenished after 12h; PV containing supernatants were collected 72h later and passed 267 through a 0.45µm filter. Transfections were carried out in 6-well Nunclon© plates (Thermo Fisher 268 Scientific, UK).

GFP-expressing pseudotypes were produced by substituting the pCSFLW firefly luciferase lentiviral vector mentioned previously with the GFP expressing vector pCSGW. PV-containing supernatants were titrated down clear 96-well Nunclon© plates (Thermo Fisher Scientific, UK) with the addition of 1x10⁴ cells per well. Plates were incubated for 72h and visualized by Nikon Eclipse 50i epiflourescence microscope with a charge-coupled QICAM Fast 1394 camera (QImaging) at x200 magnification.

275 Screening of cell lines

Two-fold serial dilutions of PV-containing supernatant were performed as previously described using white 96-well Nunclon© plates (Thermo Fisher Scientific, UK). Subsequently, approximately 1×10^4 of each cell line was added in 50µl of medium per well. Plates were incubated in a humidified incubator at 37°C and 5% CO₂ for 48h, after which 50µl of Bright-Glo[™] substrate was added and luciferase reading recorded in relative luminescence units (RLU), following a 5 minute incubation period. Control wells were used to measure cell populations for each cell line.

Pseudotype based microneutralization assay (pMN) using H17N10 and H17 bearing lentiviral pseudotypes:

284 Monoclonal antibodies (mAbs) were serially diluted 1:2 across white 96-well Nunclon© plates 285 (Thermo Fisher Scientific, UK) in 50 μ l of DMEM. PV-containing supernatants were diluted and added 286 to each well to give an approximate RLU value of 1×10^6 per well in 50 μ l of DMEM. After 1h incubation at 37°C and 5% CO₂, approximately 1×10^4 cells were added per well in a volume of 50 μ L 287 Plates were incubated for 48h at 37°C and 5% CO₂, then 50 µl of Bright-Glo[™] was added and 288 289 luminescence read after a 5 minute incubation at room temperature. Results were analyzed with 290 Graphpad (Prism 7), using nonlinear regression on luminescence values normalized to cell only and 291 virus only control thresholds (100% and 0% neutralization equivalent, respectively). IC_{50} values 292 represent the concentration (ng/ml) required for each antibody to neutralize 50% of functional 293 pseudotyped virus, based on luciferase activity.

294 Cell treatment

295 RIE1495 cells Overnight treatment of with the endosomal acidification inhibitor 296 ammonium chloride (SIGMA) and pretreatment for 2h with neuraminidase (Roche) was done as 297 previously described (Maruyama et.al, 2016). Treated cells were washed three times with serum 298 free medium, and then incubated for 24h with H17 PV. PV titration and luciferase activity was 299 monitored with a luminometer as described previously.

300 Pseudotype based enzyme linked lectin assay (pELLA) utilizing N10:

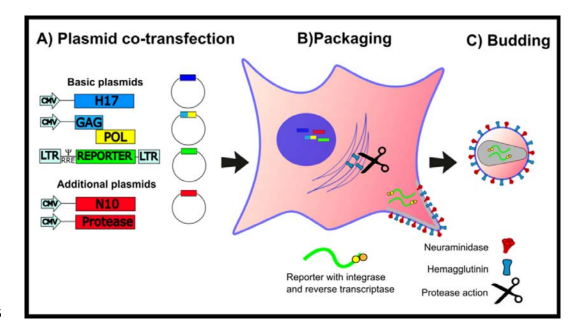
301 ELLA was performed as described by Couzens et al. 2014, but adapted in order to allow use of 302 lentiviral pseudotypes as a source of NA (Biuso et al. 2018; Prevato et al. 2015). Briefly, PV 303 containing supernatant was serially diluted (1:2) across a standard clear microtitre plate in sample 304 diluent (PBS, 1% BSA, 0.5% Tween 20). 50 μ l of the resulting dilution series was transferred in 305 duplicate to Nunclon[©] Maxisorp 96-well plates (Thermo Fisher Scientific, UK) previously coated with 306 Fetuin (Sigma Aldrich, UK) and containing 50 μ l of sample diluent per well. Plates were then 307 incubated for 18h and washed, prior to addition of conjugate diluent (PBS, 1% BSA) containing a 308 1:500 concentration of peanut lectin conjugated to horseradish peroxidase (PNA-HRPO, Sigma 309 Aldrich, UK). Plates were then incubated for 2h in the dark, whereupon OPD-based substrate in 310 citrate buffer was added (Sigma Aldrich, UK). Reactions were stopped after 10 minutes using 50 μ l 311 1M H_2SO_4 , and readings recorded using a standard ELISA plate reader at 492nm. Exogenous 312 neuraminidase from Clostridium perfringens (Sigma Aldrich, UK) was used as a positive benchmark 313 control and samples were assayed alongside PV bearing neuraminidases from other influenza 314 subtypes.

315 Statistical analysis

316 Where possible, statistical analysis was carried out to determine whether differences in PV titre 317 were significant. One-way ANOVA t-tests were performed using fold change scores with a Tukey's 318 multiple comparisons test. P-values were set at 0.05 (P[0.05) unless indicated otherwise. 319 Significantly different data are denoted with asteriks representing p= <0.01 (*), <0.001 (**), <0.0001 320 (***) and <0.00001 (****).

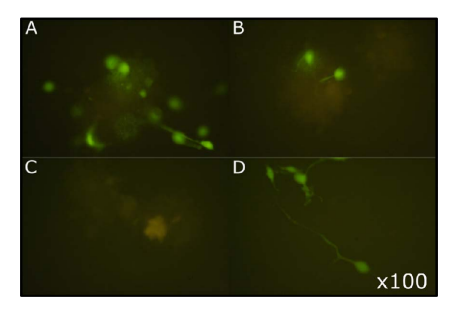
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324 Figures and Tables

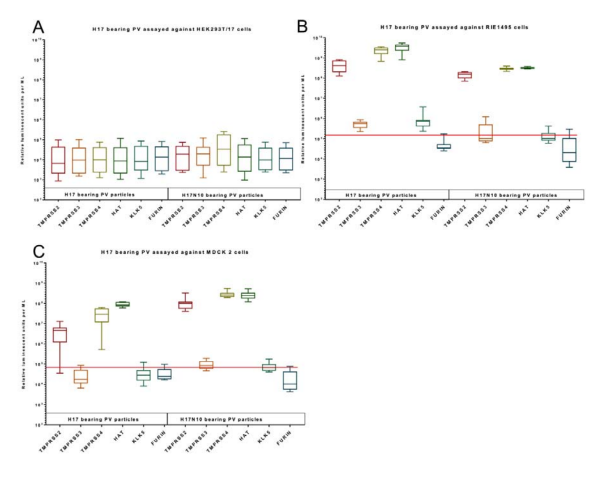


327 Figure 1. Cartoon showing the production of H17N10 PV via plasmid transfection.

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- 330 Figure 2. Transduction of U87 MG cells by influenza GFP PV (x100 magnification). A) H17 produced
- using pCAGGS-HAT. B) H17 PV produced using pCAGGS-TMPRSS2. C) Cell only control. D) H5 (A/Viet
- nam/1194/2004) PV positive control.



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Figure 3. Transduction of various target cell lines. A) HEK293T/17, B) RIE1495 and C) MDCK II with H17 and H17N10 pseudotyped viruses carrying the luciferase reporter gene. Results given in Relative

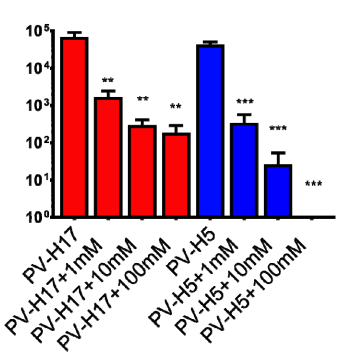
Luminescence Units per ml; RLU/ml). Average cell only luminescence shown as a red line.

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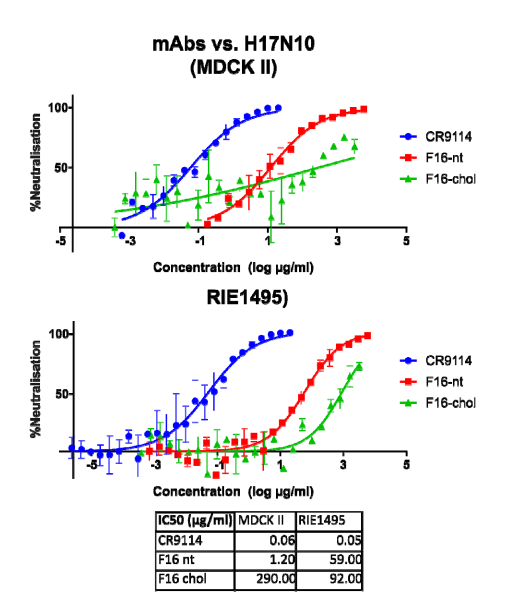
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Figure 4. The effect of ammonium chloride based acidification on the transduction of RIE1495 cells by H17 PV. Increases in acidification of RIE1495 cell cultures prior to transduction by H17 resulted in lower luciferase based titres, indicating that the pH sensitivity of H17 is similar to that of H5. Significant differences in transduction denoted by asteriks, ** represents p= <0.001 and *** p= <0.0001 respectively.



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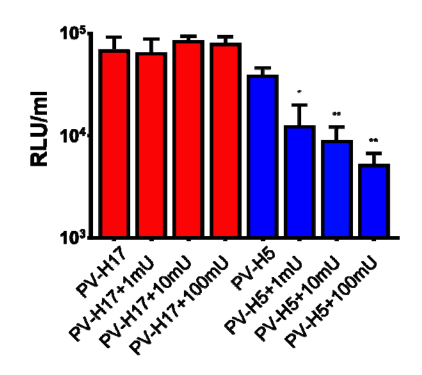
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Figure 5. Neutralization curves and IC50 values for mAbs CR9114, FI6-nt and FI6-Chol against H17N10 bearing lentiviral pseudotypes on cell lines RIE1495 and MDCK II. Nonlinear regression carried out using Graphpad (Prism 7) in order to provide IC_{50} values for each graph. Each IC_{50} is the concentration of mAb required for reduction of 50% of the virus input in terms of luciferase activity.

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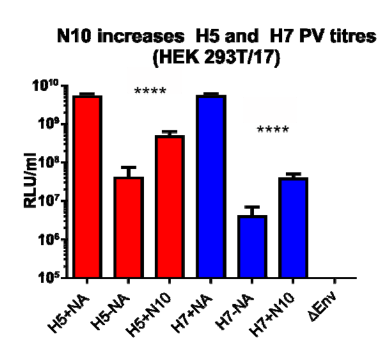


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Figure 6. The effect of neuraminidase pre-treatment on the transduction of RIE1495 cells by H17 and H5 PV. No relationship is seen between pre-treatment of cells with neuraminidase for H17 bearing PV, but H5-based transduction is significantly reduced when sialic acids are stripped from target cells. Significant difference in transduction denoted by asteriks, * represents p= <0.01 and ** p= <0.001 respectively.

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Figure 7. Release of H5 and H7 bearing PV by N10. PV bearing the glycoproteins H5 (A/Vietnam/1194/2004) and H7 (A/Shanghai/1/2013) were produced with exogenous neuraminidase (+NA), with no NA (-NA) or accompanied with an N10 glycoprotein (+N10). H5 and H7 PVs produced in the presence of N10 have significantly higher luficerase titres than those produced with no NA. Background luciferase titre control (Δ Env) control shown. Significant differences in transduction denoted by asteriks, **** represents p= <0.00001.

378 Table 1. Cell lines used in the production of Bat influenza VSV pseudotypes.

Cell Line	Species	Tissue:
Vero E6	Chlorocebus sp.	Kidney
НЕК293	Homo sapiens	Kidney
МДСК	Canis lupus familiaris	Kidney
SK-L	Sus scrofa	Kidney
QT6	Coturnix japonica	Muscle
Cell Line:	Bat species:	Tissue:
BKT1	Rhinolophus ferrumequinum	Kidney
FBKT1	Pteropus dasymallus yayeyamae	Kidney
YubFKT1	Miniopterus fuliginosus	Kidney
IndFSPT1	Pteropus giganteus	Spleen
DemKT1	Rousettus leschenaultii	Kidney
ZFBK11-97	Epomophorus gambianus	Kidney
SuBK12-08	Miniopterus schreibersii	Kidney
ZFBS13-75A	Eidolon helvum	Spleen
List of cell lines used in V	SV based pseudotyping of H17 and H18 vi	ruses. Bold entries are thos

cell lines which were found to be permissive for viral transduction by Maruyama et al. 2016.

380 Table 2: MDCK cell lines.

NBL-2 (ATCC [®] cat # CCL-34 [™]) MDCK I (EEACC cat # 00062106) MDCK.1 (EEACC cat # 00062107) MDCK.1 (ATCC [®] cat # CRL-2935 [™]) MDCK.2 (ATCC [®] cat # CRL-2286 [™]) super dome (ATCC [®] cat # CRL-2285 [™]) Different MDCK cell lines available commercially	
MDCK II (EEACC cat# 00062107) MDCK.1 (ATCC® cat# CRL-2935™) MDCK.2 (ATCC® cat# CRL- CRL-2936™) super dome (ATCC [°] cat# CRL-2286 [™]) super tube (ATCC [®] cat# CRL-2285™)	
MDCK.1 (ATCC [®] cat# CRL-2935 [™]) MDCK.2 (ATCC [®] cat# CRL- CRL-2936 [™]) super dome (ATCC [®] cat# CRL-2286 [™]) super tube (ATCC [®] cat# CRL-2285 [™])	
MDCK.2 (ATCC® cat# CRL- CRL-2936™) super dome (ATCC [®] cat# CRL-2286 [™]) super tube (ATCC® cat# CRL-2285™)	
super dome (ATCC [®] cat#CRL-2286 [™]) super tube (ATCC [®] cat# CRL-2285™)	
super tube (ATCC® cat# CRL-2285™)	
	• • •
Different MDCK cell lines available commercially	super tube (ATCC® cat# CRL-2285™)
	Different MDCK cell lines available commercially
	ple 3: Genes, plasmids and sources.

Gene	Plasmid	Source
Transmembrane protease, serine 2	pCAGGS-TMPRSS2	(Böttcher et al. 2006)
Transmembrane protease, serine 3	pcDNA3.1-hTMPRSS3	(Bertram et al. 2010)
Transmembrane protease, serine 4	phCMV-Tag3 (TMPRSS4-myc)	(Jung et al. 2008)
Human Airway Trypsin	pCAGGS-HAT	(Böttcher et al. 2006)
Kallikrein-related peptidase 5	pl.18-KLK5	Synthesised, Sigma
Furin	pFurin	Kind gift from H. D. Klenk
H17 (A/little yellow shouldered	pl.18-H17	Synthesised, Genscript
bat/Guatamala/060/2010)		
N10 (A/little yellow shouldered	pl.18-N10	Synthesised, Genscript
bat/Guatamala/060/2010)		
H5 (A/Viet Nam/1194/2004)	pl.18-H5	(Temperton et al. 2007)
HIV-1 packaging plasmid	pCMVΔR8.91 (p8.91)	(Zufferey et al. 1997)
Lentiviral vector for expression of	pHR-SIN-SE (pCSFLW)	(Demaison et al. 2002)
Firefly Luciferase		
Vesicular Stomatitis Virus	pMD.G (VSV-G)	Yasu Takeuchi, University
glycoprotein		College London

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391 Acknowledgements

EG is supported by the Biotechnology and Biological Sciences Research Council
 (http://www.bbsrc.ac.uk) via Strategic LoLa grant BB/K002465/1 "Developing Rapid Responses to
 Emerging Virus Infections of Poultry (DRREVIP)".

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