Swine ANP32A supports avian influenza virus polymerase

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Avian influenza viruses occasionally infect and adapt to mammals, including humans. Swine are often described as ‘mixing vessels’, being susceptible to both avian and human origin viruses, which allows the emergence of novel reassortants, such as the precursor to the 2009 H1N1 pandemic. ANP32 proteins are host factors that act as influenza virus polymerase cofactors. In this study we describe how swine ANP32A, uniquely among the mammalian ANP32 proteins tested, supports activity of avian origin influenza virus polymerases, and avian influenza virus replication. We further show that after the swine-origin influenza virus emerged in humans and caused the 2009 pandemic it evolved polymerase gene mutations that enabled it to more efficiently use human ANP32 proteins. We map the enhanced pro-viral activity of swine ANP32A to a pair of amino acids, 106 and 156, in the leucine-rich repeat and central domains and show these mutations enhance binding to influenza virus trimeric polymerase. These findings help elucidate the molecular basis for the ‘mixing vessel’ trait of swine and further our understanding of the evolution and ecology of viruses in this host.

Importance (max 150 words)

Avian influenza viruses can jump from wild birds and poultry into mammalian species such as humans or swine, but only continue to transmit if they accumulate mammalian adapting mutations. Pigs appear uniquely susceptible to both avian and human strains of influenza and are often described as virus ‘mixing vessels’. In this study, we describe how a host factor responsible for regulating virus replication, ANP32A, is different between swine and humans. Swine ANP32A allows a greater range of influenza viruses, specifically those from birds, to replicate. It does this through binding the virus polymerase more tightly than
the human version of the protein. This work helps to explain the unique properties of swine as 'mixing vessels'.

Introduction

Influenza A viruses continuously circulate in their natural reservoir of wild aquatic and sea birds. Occasionally, avian influenza viruses infect mammalian hosts, but these zoonotic viruses have to adapt for efficient replication and further transmission. This limits the emergence of novel endemic strains. Avian-origin, mammalian-adapted influenza viruses have been isolated from a range of mammalian species including humans, swine, horses, dogs, seals, and bats (1-6).

One mammalian influenza host of significance are swine, which have been described as susceptible to viruses of both human- and avian-origin (6). It has been hypothesised that swine act as 'mixing vessels', allowing efficient gene transfer between avian- and mammalian-adapted viruses. This leads to reassortants, which are able to replicate in humans, but to which populations have no protective antibody responses, as best illustrated by the 2009 H1N1 pandemic (pH1N1) (7). The ability of pigs to act as 'mixing vessels' has generally been attributed to the diversity of sialic acids, the receptors for influenza, found in pigs that would enable co-infection of a single host by diverse influenza strains (8, 9). The husbandry of swine has also been hypothesised to play a role in this 'mixing vessel' trait; swine are often exposed to wild birds and it is likely their environments are often contaminated with wild bird droppings containing avian influenza viruses (10, 11).

For an avian-origin influenza virus to efficiently infect and transmit between mammals several host barriers must be overcome. One major barrier is the weak activity of avian influenza virus polymerases in the mammalian cell (12, 13). The acidic, (leucine-rich)
nuclear phosphoproteins of 32 kilodaltons (ANP32) proteins are key host factors responsible for the restricted polymerase activity of avian influenza viruses in mammalian cells (14). ANP32 proteins possess an N-terminal domain composed of five leucine rich repeats (LRRs) and a C-terminal low complexity acidic region (LCAR) separated by a short region termed the ‘central domain’. In birds and most mammals three ANP32 paralogues are found: ANP32A, ANP32B and ANP32E (15, 16). The roles of ANP32 proteins in cells are diverse and often redundant between the family members but include histone chaperoning, transcriptional regulation, regulation of nuclear export and apoptosis (16). In birds, such as chickens and ducks, an exon duplication allows for the expression of an alternatively spliced, longer isoform of ANP32A that effectively supports activity of polymerases of avian influenza viruses (14, 17). Mammals only express the shorter forms of ANP32 proteins which do not efficiently support avian polymerase unless the virus acquires adaptive mutations, particularly in the PB2 polymerase subunit, such as E627K (14). A further difference between the ANP32 proteins of different species is the level of redundancy in their ability to support influenza polymerase. In humans, two paralogues – ANP32A and ANP32B – are essential but redundant influenza polymerase cofactors (18, 19). In birds, only a single family member – ANP32A - supports influenza virus polymerase activity, as avian ANP32B proteins are not orthologous to mammalian ANP32B (15, 19, 20). In mice, only ANP32B can support influenza A polymerase activity (18, 19). Neither avian nor mammalian ANP32E proteins have been shown to support influenza polymerase activity (18-20).

In this study, we investigated the ability of a variety of mammalian ANP32 proteins to support influenza virus polymerases derived from viruses isolated from a range of hosts. We find differences in pro-viral efficiency that do not always coincide with the natural virus-host relationship: for example, human ANP32B is better able to support bat influenza
polymerases than either bat ANP32 protein. Conversely, we describe evidence of human ANP32 adaptation early during the emergence of the pH1N1 virus from pigs, and find that swine ANP32A is the most potent pro-viral mammalian ANP32 protein tested, supporting non-adapted avian virus polymerase activity and avian influenza virus replication significantly better than human ANP32A. This can be attributed to amino acid differences in the LRR4 and central domains that enhance the interaction between swine ANP32A and the influenza polymerase complex, suggesting a mechanism for this enhanced pro-viral activity. Our findings give support to the special status as potential ‘mixing vessels’ of swine in influenza evolution.

Results

Mammals naturally susceptible to influenza have two pro-viral ANP32 proteins.

To investigate the ability of different mammalian ANP32A and ANP32B proteins to support influenza virus polymerase activity, several mammalian-origin influenza virus polymerase constellations were tested using an ANP32 reconstitution minigenome assay. A previously described human cell line with both ANP32A and ANP32B ablated (eHAP dKO) was transfected with expression plasmids encoding ANP32A or ANP32B from chicken, human, swine, horse, dog, seal or bat, as well as the minimal set of influenza polymerase expression plasmids for PB2, PB1, PA and nucleoprotein (NP), to drive amplification and expression of a firefly-luciferase viral-like reporter RNA and a Renilla-luciferase expression plasmid as a transfection control.

Initially, we tested a panel of polymerases derived from human, canine, equine and bat influenza viruses. In contrast to chicken ANP32B, which does not support influenza virus polymerase activity (15, 19, 20), chicken ANP32A and all mammalian ANP32A and ANP32B
proteins supported activity of the mammalian-origin viral polymerases to varying degrees 
(Fig. 1a). Among the mammalian ANP32 proteins tested, for most polymerases, swine 
ANP32A provided the strongest support of polymerase activity, whereas the ANP32B 
proteins from dog, seal and bat displayed the least efficient pro-viral activity, lower than 
those species’ respective ANP32A proteins. These trends could not be explained by 
differences in expression levels or nuclear localisation (Fig. 1b, c). The bat influenza 
polymerases, along with (human) influenza B polymerase showed a different pattern of 
ANP32 usage, being able to strongly utilise ANP32Bs from all mammalian species, 
particularly human ANP32B (Fig. 1a). There was no evidence that influenza viruses adapted 
to particular mammals had evolved to specifically use the corresponding ANP32 proteins. 
For example, dog ANP32A or ANP32B were not the most efficient cofactors for canine 
influenza virus polymerase and human ANP32B was better able to support the bat influenza 
polymerase than either of the bat ANP32 proteins.

Swine ANP32A, but not other mammalian ANP32 proteins, can support the polymerase 
activity and virus replication of avian-origin influenza viruses

We next tested the ANP32 preference of a human 2009 (swine-origin) pH1N1 and 
two polymerases from swine influenza isolates. Interestingly, these polymerases were 
robustly supported by chicken and swine ANP32A, but not other mammalian ANP32 
proteins, with the Eurasian avian-like polymerase from A/swine/England/453/2006 (H1N1; 
sw/453) showing the clearest effect (Fig. 2a). We went on to test a panel of avian-origin viral 
polymerases with no known mammalian polymerase adaptations, including 
A/duck/Bavaria/77(H1N1; Bav), thought to be an avian precursor of the Eurasian avian-like 
swine lineage (Fig. 2b)(5). For all the avian origin viral polymerases the stringent preference
for avian ANP32A to support polymerase activity was evident (co-expression of chicken ANP32A led to very strong polymerase activity). However, amongst all the mammalian ANP32 proteins tested, only swine ANP32A was able to significantly support avian influenza polymerase activity, though to a lesser degree than chicken ANP32A (Fig. 2b). This unique pro-viral effect of swine ANP32A on swine and avian-origin polymerases was maintained across a wide titration of plasmid doses (Fig. 2c).

Furthermore, we tested the relative ability of human and swine cells to support replication of a non-adapted avian influenza virus. Isogenic recombinant A/turkey/England/50-92/1991(H5N1; 50-92) virus containing either wild-type PB2 (E627) or the mammalian adaptation PB2-E627K were used to infect wild type human eHAP and swine NPTr cells (Fig 3a). Although E627K significantly increased the virus replication in both cell lines, the magnitude of difference was less in the swine cells than the human cells at earlier time points (for example 17-fold vs 110x-fold at 12 hours post-infection). The less drastic reduction in replication of the virus with non-adapted avian origin polymerase compared with the adapted control in swine cells is consistent with the hypothesis that swine ANP32A can support replication of avian influenza viruses.

To investigate whether this difference was indeed accounted for by differences in ANP32A proteins, chicken, swine, or human ANP32A were pre-expressed in eHAP dKO cells that were then infected with 50-92 wild type and E627K recombinant viruses (Fig. 3b). As shown previously, when empty vector was expressed no virus replication took place (18). For the mammalian adapted PB2-E627K virus it made little difference which ANP32A protein was expressed although a trend was seen for chicken ANP32A supporting higher titres than swine ANP32A, which, in turn, supported higher titres than human ANP32A. For the non-
adapted PB2-E627 virus, however, a greater and significant difference was seen – chicken ANP32A clearly supported virus replication better than either mammalian ANP32A protein. Swine ANP32A supported replication of the avian influenza virus to a higher level than human ANP32A at all time points, and this difference was significant (P<0.05) at 24 hours post infection. Overall, this indicates that swine ANP32A is better able to both support avian influenza virus polymerase activity, as well as virus replication, than human ANP32A.

The pH1N1 swine influenza virus polymerase, adapting to humans, evolved to better use human ANP32 proteins

In 2009 the swine-origin pH1N1 influenza virus adapted from pigs for transmission between humans causing an influenza pandemic (7). The pH1N1 polymerase genes were derived from a swine triple reassortant constellation in which PB2 and PA originally derived from avian influenza viruses in the mid-1990s (21). From 2009 to 2010 the virus continued to circulate and adapt to humans in the second and third pandemic waves (22). pH1N1 viruses contain the PB2 polymerase adaptations T271A, G590S, and Q591R, which appear to compensate for the lack of E627K in enabling replication in mammalian cells and these amino acids did not change between the first and third waves of the pandemic (23). We previously showed that a single substitution in the PA subunit of the polymerase, N321K, contributed to increased polymerase activity of third-wave pH1N1 viruses in human cells (22). We hypothesised that this PA mutation might function by improving support for the emerging virus polymerase by the human ANP32 proteins.

We performed minigenome assays with a first-wave pandemic virus, A/England/195/2009(pH1N1; E195), and a third-wave pandemic virus A/England/687/2010(pH1N1; E687), which differ in PA at position 321. As shown before, PA
321K enhances polymerase activity in general in both virus polymerase backgrounds in human eHAP cells, as well as swine NPr cells (Fig. 4a). However, the boost is far greater in the human cells (~7-fold) than in the swine cells (~2-fold), implying this mutation may have arisen to overcome the greater restriction seen upon the jump into humans (22).

We next tested the ability of human and swine ANP32 proteins to support the different pH1N1 polymerases in eHAP dKO cells. Polymerases containing PA-321N are more robustly enhanced by swine ANP32A (by around 3.5-fold compared to human ANP32A), as is typical of swine-origin polymerases (Fig. 4b). Swine ANP32A, however, gives a much more modest boost to polymerase activity compared to human ANP32A when 321K is present (<2-fold). This suggests the PA N321K adaptation was selected in these viruses to adapt to the more poorly supportive ANP32 proteins present in human cells. We could further show that endogenous swine ANP32A protein is predominantly localised in the nucleus in swine NPr cells, consistent with our previous over-expression data (Fig. 4c).

Differences in swine and human ANP32A pro-viral activity can be mapped to the LRR4 and central region.

We set out to identify the molecular basis for the unusually high activity of swine ANP32A in comparison with the other mammalian ANP32 proteins. An alignment of ANP32A primary sequences identified three amino acids outside the LCAR, that differed between swine ANP32A and the other mammalian orthologues. Using reciprocal mutant ANP32A proteins, the identity of amino acid position 156, naturally a serine in swine ANP32A but a proline in most other mammalian and all avian ANP32A proteins, was shown to have a major, reciprocal influence on activity (Fig. 5a). The amino acid at position 106 contributed to a lesser degree, with swine-like valine enhancing pro-viral activity over human-like
isoleucine when complementing the swine influenza polymerase constellation, though changes at this residue appeared to have more minor effects on proviral activity supporting the 50-92 and Bav avian virus polymerases. Position 228, localised nearby the C-terminal nuclear localisation signal of ANP32A, had no appreciable impact. In the background of human ANP32A, I106V generally gave between a 1.5- and 6-fold increase in polymerase activity while P156S gave between a 3- and 16-fold boost, depending on the polymerase constellation tested. The combined 106/156 mutant showed an additive effect implying these two residues are, together, responsible for the enhanced pro-viral activity of swine ANP32A (Fig. 5a,c). None of the mutations affected expression levels (Fig. 5b). Positions 106 and 156 map to the LRR4 and central domains of ANP32 protein, respectively, proximal to the previously characterised LRR5 residues, 129/130, that are responsible for the lack of pro-viral activity of avian ANP32B proteins (Fig. 5c)(15, 19). This reinforces the concept that the LRR4/LRR5/central region of ANP32 proteins is essential to their pro-viral function. Indeed, we could show that introducing the mutation N129I into swine ANP32A abrogated its ability to support influenza polymerase activity (Fig. 5a).

An increase in binding to the polymerase accounts for the enhanced pro-viral activity of swine ANP32A

Pro-viral ANP32 proteins from birds and mammals directly bind trimeric polymerase in the cell nucleus (17, 24, 25). Moreover, the inability of avian ANP32B to support influenza polymerase activity correlates with a lack of protein interaction conferred by amino acid differences at residues 129 and 130 (15).

To assess the strength of interaction between swine ANP32A protein and influenza polymerase, we used a split-luciferase assay, where the two halves of Gaussia luciferase are
fused onto PB1 and ANP32 protein (15, 25). As seen previously (25), the interaction
between influenza virus polymerase and human ANP32A was weak but detectable above
background (huA, Fig. 6a). Swine ANP32A interacted more strongly with both human-origin -
E195 (pH1N1 2009) - and avian-origin - A/turkey/England/50-92/1991(H5N1) - influenza
polymerases, although not as strongly as chicken ANP32A (Fig. 6a). Furthermore, the two
residues identified as being responsible for strong pro-viral activity of swine ANP32A, at
positions 106 and 156, enhanced polymerase binding by human ANP32A and the reciprocal
mutations decrease the swine ANP32A interaction, implying the mode of action of these
mutations is through enhancing swine ANP32A-polymerase interactions (Fig. 6a). It was also
shown that N129I, the substitution naturally identified in chicken ANP32B and previously
shown to abolish binding and activity in chicken and human ANP32 proteins (15, 19),
showed a similar phenotype in swine ANP32A, abolishing detectable binding and activity
(Fig. 6a,b). The ablations of the pro-viral activity of swine ANP32A and ANP32B by the
substitution N129I was not explained by reductions in expression of these mutant proteins
(Fig. 6b,c).

Estimating the pro-viral activity of ANP32A proteins from other mammalian species

Based on the molecular markers described in this study it is possible to survey
ANP32A proteins from all mammals to predict which other species may have highly
influenza polymerase supportive proteins and therefore potential to act as mixing vessels
for reassortment between avian and mammalian-adapted influenza viruses.

Very few mammals share the pro-viral marker, 156S, and the few that do mostly
constitute species not yet described as hosts for influenza viruses (Fig. 6d). A notable
exception is the pika which, in a similar manner to pigs, are known to often become infected
with avian influenza viruses with minimal mammalian adaptation (26-28). Pigs are currently the only known mammalian species with a publicly available ANP32A sequence that contain the secondary, minor pro-viral maker 106V.

Discussion

In this study we describe the ability of different mammalian ANP32A and ANP32B proteins to support activity of influenza virus polymerases isolated from a variety of hosts. We found that swine ANP32A, uniquely among the ANP32 proteins, supports avian influenza virus polymerase activity and virus replication. Swine ANP32A does not harbour the avian-specific 33 amino acid duplication that enables the strong interaction and efficient support of polymerase activity of avian-origin viruses by avian ANP32A proteins (14). Thus, avian influenza viruses are restricted for replication in swine as we have previously shown, and mammalian-adapting mutations enhance their polymerase activity in pig cells (11). Nonetheless, this level of pro-viral activity associated with swine ANP32A, albeit weaker than avian ANP32As, may contribute to the role of swine as mixing vessels: non-adapted avian influenza viruses that infect pigs could replicate sufficiently to accumulate further mutations that allow for more efficient mammalian adaptation and/or reassortment, enabling virus to either become endemic in swine or to jump into other mammals, including humans.

We map this strongly pro-viral polymerase phenotype to a pair of mutations which allow swine ANP32A to bind more strongly to influenza virus polymerase, potentially explaining the mechanism behind its enhanced pro-viral activity. These residues are only found in a handful of other mammals including pika. It is conceivable these residues are
located at a binding interface between polymerase and ANP32, but resolution of the structure of the host:virus complex will be required to confirm this hypothesis.

A recent study from Zhang and colleagues independently corroborated the superior ability of swine ANP32A amongst mammalian ANP32 proteins to support avian influenza virus polymerase activity (29). Moreover, they also correlated this phenotype with amino acids at position 106 and 156 that increased the strength of interactions between the host factor and the viral polymerase complex. In their studies the interaction between ANP32 proteins and viral polymerase was measured by co-immunoprecipitation, making it unlikely that the similar differences we measured using our quantitative split luciferase assay were due to re-orientation of the luciferase tags.

It has long been speculated that swine play a role as ‘mixing vessels’, by acting as host to both human- and avian-origin influenza viruses (30). This trait may be partially attributed to receptor patterns in swine allowing viruses that bind to both α2,3 linked (i.e. avian-like viruses) and α2,6 linked sialic acid (i.e. human-like) to replicate alongside each other (8, 9). However, replication of the avian-origin influenza virus genomes inside infected cells is also required to enhance the opportunity for further adaptation or reassortment. We previously developed a minigenome assay for assessing polymerase activity in swine cells and showed that avian virus polymerases were restricted and that restriction could be overcome by typical mutations known to adapt polymerase to human cells (11). Taken together the ability to enter swine cells without receptor switching changes in the haemagglutinin gene, along with a greater mutation landscape afforded in swine cells by the partially supportive pro-viral function of swine ANP32A may have an additive effect to allow swine to act an intermediate host for influenza viruses to adapt to mammals. Furthermore,
our work implies other mammals, such as the pika, could play a similar role which is of particular interest due to the pika’s natural habitat often overlapping with that of wild birds and its (somewhat swine-like) distribution of both α2,3 and α2,6-linked sialic acid receptors (31).

Upon crossing into humans from swine, it is likely that viruses would be under selective pressure to adapt to human pro-viral factors, such as the ANP32 proteins. We use the example of a pair of first- and third-wave pandemic H1N1 influenza viruses isolated from clinical cases in 2009 and 2010 (22). The polymerase constellation of the 2009 pH1N1 virus contains PB2 and PA gene segments donated from avian sources to a swine virus in a triple reassortant constellation in the mid-1990s, then passed onto humans in 2009 (21). Although the first-wave viruses, derived directly from swine, can clearly replicate and transmit between humans, over time the PA substitution, N321K, was selected because it enabled more efficient activity of the viral polymerase in human cells. Our data suggests this is a direct adaptation to human ANP32 proteins. This again illustrates how swine have acted as a ‘halfway house’ for the step-wise adaptation of genes originating in avian influenza viruses that have eventually become humanised.

Also of note, we show here that as for the human orthologues (18, 19), the ANP32A and B proteins of swine (as well as all other mammals tested here) are redundant in their ability to support the viral polymerase. We further show that the substitution N129I is able to partially or fully ablate the pro-viral activity of swine ANP32A and ANP32B. We suggest that the introduction of this substitution in both swine ANP32A and ANP32B by genome editing would be a feasible basis for generating influenza resistant, or resilient, pigs, in a
similar manner to that demonstrated for porcine respiratory and reproductive syndrome virus resistant pigs, and proposed for influenza resistant, or resilient, chickens (15, 32).

To conclude, we hypothesise that the superior pro-viral function of swine ANP32A for supporting influenza replication may enable swine to act as intermediary hosts for avian influenza viruses, and also affect the way the viruses evolve as they pass from birds, through swine, and onto humans. This, in turn, may influence the ability of different swine influenza viruses to act as zoonotic agents or as potential pandemic viruses.

Materials and methods

Cells

Human engineered-Haploid cells (eHAP; Horizon Discovery) and eHAP cells with ANP32A and ANP32B knocked out (dKO) by CRISPR-Cas9, as originally described in (18), were maintained in Iscove’s Modified Dulbecco’s Medium (IMDM; ThermoFisher) supplemented with 10% fetal bovine serum (FBS; Biosera), 1% non-essential amino acids (NEAA; Gibco) and 1% Penicillin-streptomycin (pen-strep; invitrogen). Human embryonic kidney (293Ts, ATCC), Newborn Pig Trachea cells (NPtr; ATCC), and Madin-Darby Canine Kidney cells (MDCK; ATCC) were maintained in Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 10% FBS, 1% NEAA and 1% pen-strep. All cells were maintained at 37°C, 5% CO₂.

ANP32 plasmids constructs

Animal ANP32 constructs were codon optimised and synthesised by GeneArt (ThermoFisher). Sequences used were pig (Sus scrofa) ANP32B (XP_020922136.1), Horse (Equus caballus) ANP32A (XP_001495860.2) and ANP32B (XP_023485491.1), Dog (Canis
lupus familiaris) ANP32A (NP_001003013.2), Dingo (Canis lupus dingo) ANP32B (XP_025328134.1), Monk Seal (Neomonachus schauinslandi) ANP32A (XP_021549451.1) and ANP32B (XP_021546921.1), and Common Vampire Bat (Desmodus rotundus) ANP32A (XP_024423449.1) and ANP32B (XP_024415874.1). All isoforms were chosen based on their orthology and synteny to the known functional human isoforms. Species of origin were chosen due to being influenza hosts or the most-commonly related species to influenza hosts (in the case of Monk Seal which are closely related to Harbour Seal whereas common vampire bats belong to the same family as little yellow-shouldered and flat-faced bats). Dingo ANP32B was substituted for dog ANP32B as the equivalent isoform used for all other ANP32Bs is unannotated in the dog genome due to a gap in the scaffold. All ANP32 expression constructs included a C-terminal GSG-linker followed by a FLAG tag and a pair of stop codons. Overlap extension PCR was used to introduce mutations into the ANP32 constructs which were then subcloned back into pCAGGS and confirmed by Sanger sequencing.

Viral minigenome plasmid constructs

Viruses and virus minigenome full strain names used through this study were:

- A/Victoria/1975(H3N2; Victoria), A/England/195/2009(pH1N1; E195),
- A/England/687/2010(pH1N1; E687), A/Japan/WRAIR1059P/2008(H3N2; Japan),
- B/Florida/4/2006 (B/Florida), A/Anhui/2013(H7N9; Anhui), A/duck/Bavaria/1/1977(H1N1, Bavaria), A/turkey/England/50-92/1991(H5N1; 50-92), A/chicken/Pakistan/UDL-01/2008(H9N2; UDL1/08), A/canine/New York/dog23/2009(H3N8; CIV H3N8),
- A/canine/Illinois/41915/2015(H3N2; CIV H3N2), A/equine/Richmond/1/2007(H3N8; Richmond), A/swine/England/453/2006(EAH1N1; sw/453), A/swine/Hubei/221/2016(H1N1;
Hubei), A/little yellow-shouldered bat/Guatemala/164/2009(H17N10; H17) and A/flat-faced bat/Peru/033/2010(H18N11; H18). Viral minigenome expression plasmids (for PB2, PB1, PA and NP) for H3N2 Victoria, H5N1 50-92, H1N1 E195, H1N1 E687 IBV Florida/06, H9N2 UDL1/08 and H1N1 Bavaria have been previously described (11, 14, 22, 33). Viral minigenome plasmids for H1N1 swine/453, H3N2 Japan, H3N2 CIV, H3N8 CIV, Hubei and Richmond were subcloned from reverse genetics plasmids or cDNA into pCAGGS expression vectors using virus segment specific primers.

pCAGGs minigenome reporters for H17N10 and H18N11 bat influenza viruses were a kind gift from Professor Martin Schwemmle, Universitätsklinikum Freiburg (34). pCAGGs minigenome reporters for H7N9 were a kind gift from Professor Munir Iqbal, The Pirbright Institute, UK. Reverse genetics plasmids for H3N8, Richmond were a kind gift from Adam Rash of the Animal Health Trust, Newmarket, UK. Reverse genetics plasmids for H3N2 CIV and H3N8 CIV were a kind gift from Dr. Colin Parrish of the Baker Institute for Animal Health, Cornell University (35, 36). Viral RNA from sw/453 was kindly provided by Dr. Sharon Brookes, Animal Plant and Health Agency, Weybridge, UK.

Minigenome assay

eHAP dKO cells were transfected in 24 well plates using lipofectamine® 3000 (thermo fisher) with a mixture of plasmids; 100ng of pCAGGs ANP32/pCAGGs empty, 40ng of pCAGGs PB2, 40ng of pCAGGs PB1, 20ng of pCAGGs PA, 80ng of pCAGGs NP, 40ng of pCAGGs Renilla luciferase, 40ng of poll vRNA-Firefly luciferase. Transfections in wild-type eHap cells were performed similarly but without ANP32. Transfections in NPTr cells were carried out in 12 well plates using the same ratios above. 24 hours post-transfection cells were lysed with passive lysis buffer (Promega) and luciferase bio-luminescent signals were...
read on a FLUOstar Omega plate reader (BMG Labtech) using the Dual-Luciferase® Reporter

Assay System (Promega). Firefly signal was divided by Renilla signal to give relative
luminescence units (RLU). All assays were performed with 2 or 3 separate repeats on
different days, representative experiments are shown.

Viruses replication assays

All virus replication assays were performed with recombinant viruses containing the
HA, NA and M genes of A/Puerto Rico/8/1934(H1N1; PR8) and the remaining genes from
the avian influenza virus 50-92 containing PB2 627 E (wild type) or K, as has been described
previously (11). eHAP dKO cells pre-transfected 24 hours prior with 400ng of pCAGGs-
ANP32A (chicken, swine or human) or pCAGGs-empty, or wild type eHAP or NPTr cells were
infected at a multiplicity of infection of 0.001 in 6 well plates. Virus growth media, either
IMDM or DMEM (for eHAP cells and NPTr cells, respectively) was made from serum-free
media containing 1 μg/ml of N-tosyl-L-phenylalanine chloromethyl ketone-treated trypsin
(Worthington-Biochemical). Virus containing supernatants were collected at 12, 24, 48, 72
hours post-inoculation and stored at -80°C. Titres were assessed by infectious plaques on
MDCKs. All time points were taken in triplicate and all virus growth curves were performed
at least twice with a representative repeat shown.

Split Luciferase Assay

Split luciferase assays were undertaken in 293Ts seeded in 24 well plates. 30ng each
of PB2, PA, and PB1, with the N-terminus of Gaussia Luciferase (Gluc1) tagged to its C-
terminus after a GGSGG linker, were co-transfected using lipofectamine 3000 along with
ANP32A, tagged with the C-terminus of Gaussia Luciferase (Gluc2) on its C-terminus (after a
GGSGG linker). 24 hours later cells were lysed in 100μl of Renilla lysis buffer (Promega) and
Gaussia activity was measured using a Renilla luciferase kit (Promega) on a FLUOstar Omega 408 plate reader (BMG Labtech). Normalised luminescence ratios (NLR) were calculated by dividing the values of the tagged PB1 and ANP32 wells by the sum of the control wells which contained 1) untagged PB1 and free Gluc1 and 2) untagged ANP32A and free Gluc2 as described elsewhere (15, 37).

**Western Blotting**

To confirm equivalent protein expressing during mini-genome assays transfected cells were lysed in RIPA buffer (150mM NaCl, 1% NP-40, 0.5% Sodium deoxycholate, 0.1% SDS, 50mM TRIS, pH 7.4) supplemented with an EDTA-free protease inhibitor cocktail tablet (Roche).

Membranes were probed with mouse α-FLAG (F1804, Sigma), rabbit α-Vinculin (AB129002, Abcam), rabbit α-PB2 (GTX125926, GeneTex) and mouse α-NP ([C43] ab128193, Abcam). The following near infra-red (NIR) fluorescent secondary antibodies were used: IRDye® 680RD Goat Anti-Rabbit (IgG) secondary antibody (Ab216777, Abcam) and IRDye® 800CW Goat Anti-Mouse (IgG) secondary antibody (Ab216772, Abcam). Western Blots were visualised using an Odyssey Imaging System (LI-COR Biosciences).

**Immunofluorescence**

For investigating localisation of exogenously expressed ANP32 proteins, eHAP ANP32 dKO cells were cultured on 8 well chambered cover slips (Ibidi) and transfected with 125 ng of the indicated FLAG-tagged ANP32 protein. Cells were fixed in PBS, 4% paraformaldehyde 24 hours post transfection, then permeabilised in PBS, 0.2% Triton X-100. Cells were blocked in PBS, 2% bovine serum albumin and 0.1% tween. FLAG-tagged ANP32 proteins were detected using mouse anti-FLAG M2 primary antibody (Sigma), followed by goat anti-mouse
Alexa Fluor 568 (Invitrogen). Nuclei were counterstained with DAPI. Images were obtained using a Zeiss Cell Observer widefield microscope with ZEN Blue software, using a Plan-Apochromat 63x 1.40-numerical aperture oil objective (Zeiss) and processed using FIJI software (38).

For investigating endogenous levels of ANP32A in swine cells, NPtr cells were cultured in Nunc™ 24 well tissue culture plates on cover slips (VWR) preincubated with 10% (v/v) collagen (Rat’s tail, Sigma-Aldrich) in PBS. Cells were fixed with PBS, 4% paraformaldehyde for 20 minutes at room temperature. Cells were permeabilized with PBS, 1% Triton X-100 for 10 minutes, followed by 3 washes with PBS 0.1% Triton X-100 and blocking with PBS, 5% (w/v) skim milk powder for 1 hour at room temperature. ANP32A was detected using ab189110 (Abcam) incubated in PBS, 5% (w/v) skim milk powder overnight at 4°C, followed by incubation with anti-rabbit AlexaFluor488 (ab150077, Abcam). Phalloidin was detected using an AlexaFluor647 conjugated antibody (ab176759, Abcam), incubated during the secondary antibody application step at 1:10,000. Nuclei were counterstained with DAPI (1:15,000, Thermo Fisher). Images were captured with a Leica DMLB fluorescence microscope using Micro-Manager software at 40x or 20x for DAPI and Phalloidin respectively. Images were processed using FIJI software.

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Figure legends

Figure 1 – Most common mammalian influenza hosts have two ANP32 proteins capable of supporting influenza polymerase. a) Minigenome assays performed in human eHAP dKO with ANP32 proteins from different avian or mammalian species co-transfected. Green bars indicate species the influenza virus polymerase was isolated from, orange bars indicate recent species the virus has jumped from. Data indicates triplicate repeats plotted as mean with standard deviation. Data for each polymerase normalised to chicken ANP32A. b) Western blot assay showing protein expression levels of FLAG-tagged ANP32 proteins, NP and PB2 during a minigenome assay. c) Immunofluorescence images showing nuclear localization of the ANP32 proteins.
localisation of all FLAG-tagged ANP32 proteins (red) tested. Nuclei are stained with DAPI (blue). Abbreviations: ch – chicken, hu – human, sw – swine, eq – equine. Statistical significance was determined by one-way ANOVA with multiple comparisons against empty vector or between ANP32 proteins from the same host. *, 0.05 ≥ P > 0.01; **, 0.01 ≥ P > 0.001; ***, 0.001 ≥ P > 0.0001; ****, P ≤ 0.0001.

**Figure 2.** swANP32A can support the activity of minimally mammalian-adapted or completely non-adapted polymerases. Minigenome assays of swine (a) and avian (b) polymerases performed in human eHAP dKO cells with ANP32 proteins from different avian or mammalian species co-transfected. Green bars indicate species the influenza virus polymerase was isolated from, orange bars indicate recent species the virus has jumped from. Data indicates triplicate repeats plotted as mean with standard deviation. Data for each polymerase normalised to chicken ANP32A. c) ANP32 protein titrations with three different virus polymerase constellations. ANP32 expression plasmids were diluted in a series of 3x dilutions starting with 100ng. Data indicates triplicate repeats plotted as mean with standard deviation. Statistical significance was determined by one-way ANOVA with multiple comparisons against empty vector. **, 0.01 ≥ P > 0.001; ***, 0.001 ≥ P > 0.0001; ****, P ≤ 0.0001.

**Figure 3.** Swine ANP32A can support avian influenza virus replication better than human ANP32A. Comparative growth kinetics of isogenic, recombinant avian influenza viruses (A/turkey/England/50-92/1991(H5N1)) PB2 627E (wild type) vs E627K in (a) wild-type human eHAP cells and swine NPTr cells and (b) eHAP dKO cells pre-expressing empty vector, chicken, swine or human ANP32A. Cells were infected at a multiplicity of infection (MOI) of
All time points taken in triplicate and mean viral titres determined by plaque assay in MDCK cells with standard deviation shown. Graph is representative data of at least two independent repeats showing the same trends. Statistical significance determined by multiple Student’s t-tests in panel (a) and one-way ANOVA with multiple comparisons in panel (b). Value shown on graph in panel (a) indicate fold-change in mean titres. Dotted lines on graphs indicate limits of detection. *, 0.05 ≥ P > 0.01; **, 0.01 ≥ P > 0.001; ***, 0.001 ≥ P > 0.0001; ****, P ≤ 0.0001.

Figure 4. Third-wave pandemic H1N1 viruses adapt to human ANP32 proteins through the PA mutation N321K. a) Minigenome assays of polymerases derived from first- and third-wave pH1N1 viruses (E195 and E687, respectively) performed in wild-type human eHAP cells and swine NPTr cells. Data indicates triplicate repeats plotted as mean with standard deviation. Data normalised to E195 wild type. b) Minigenome assays performed in human eHAP cells with ANP32A and ANP32B knocked out and complemented with ANP32 proteins from human or swine following co-transfection of expression plasmids. Data indicates triplicate repeats plotted as mean with standard deviation. Data normalised to E195 wt with chicken ANP32A. All experiments in parts a) and b) performed on two separate occasions with a representative repeat shown. c) Indirect immunofluorescence images showing endogenous nuclear localisation of swine ANP32A in swine NPTr cells. Statistical significance was determined by one-way ANOVA with multiple comparisons. ****, P ≤ 0.0001.

Figure 5. The enhanced pro-viral activity of swine ANP32A maps to amino acids in LRR4 and the central domain. a) Minigenome assays with polymerase constellations from a swine
or an avian influenza virus performed in human eHAP dKO cells with human/swine ANP32A reciprocal mutants expressed. Data indicates triplicate repeats plotted as mean with standard deviation repeated on two separate occasions with a representative repeat shown. Data normalised to each polymerase with swine ANP32A wild type. b) Western blot analysis showing expression levels of human/swine ANP32A from minigenome assays. c) Crystal structure of ANP32A (PDBID: 2JE1) with residues found to affect pro-viral activity mapped (39). The unresolved, unstructured LCAR shown as a yellow line. Schematic made using PyMol (40). Statistical significance was determined by one-way ANOVA with multiple comparisons. *, 0.05 ≥ P > 0.01; ***, 0.001 ≥ P > 0.0001; ****, P ≤ 0.0001.

**Figure 6. Amino acid residues responsible for the enhanced support of polymerase activity of swine ANP32A also mediate increased binding to influenza trimeric polymerase.** a) Split luciferase assays showing the relative binding of different ANP32 proteins to trimeric polymerase from human pH1N1 or avian H5N1 viruses. PB1 was tagged with the N-terminal part of Gaussia luciferase while ANP32 proteins were tagged with the C-terminal part. NLR, normalised luminescence ratio, calculated from the ratio between tagged and untagged ANP32/PB1 pairs. Assay performed in 293T cells. Data indicates triplicate repeats plotted as mean with standard deviation, repeated across two separate experiments with representative data shown. Statistical significance was determined by one-way ANOVA with multiple comparisons between the swA and huA wild-types and mutants. ***, 0.001 ≥ P > 0.0001; ****, P ≤ 0.0001. b) Minigenome assays with reconstituted polymerases from 3 different influenza viruses, performed in human eHAP cells with ANP32A and ANP32B knocked out and complemented with wild type swine ANP32A or B or N129I mutants thereof. Data indicates triplicate repeats plotted as mean with standard deviation, repeated
across two separate experiments with representative data shown. Data normalised to each polymerase with wild type swine ANP32A. c) Western blot assay showing protein expression levels of FLAG-tagged swine ANP32 wild type or N129I proteins during a minigenome assay. d) Phylogenetic tree of mammalian ANP32A proteins. Species which contain the highly pro-viral 156S shown in red, species with 156P shown in black. Phylogenetic trees made using the neighbour-joining method based on amino acid sequence. Statistical significance was determined by one-way ANOVA with multiple comparisons against empty vector. ****, P ≤ 0.0001.
