Influenza A virus infection of primary differentiated airway epithelial cell cultures derived from Syrian golden hamsters

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Abstract

The ability of several different influenza A virus strains to infect and replicate in primary, differentiated airway epithelial cell cultures from Syrian golden hamsters was investigated. All virus strains tested replicated equivalently in the cultures and displayed a preference for infecting nonciliated cells. This tropism correlated with the expression of both α2,3- and α2,6-linked sialic acid on the nonciliated cells. In contrast, the ciliated cells did not have detectable α2,6-linked sialic acid and expressed only low amounts of α2,3-linked sialic acid. In contrast to clinical isolates, laboratory strains of influenza A virus infected a limited number of ciliated cells at late times post-infection. The presence of α2,3- and α2,6-linked sialic acid residues on the same cell type suggests that Syrian golden hamsters and differentiated airway epithelial cell cultures derived from hamsters may provide a system for studying the reassortment of influenza A virus strains which utilize different forms of sialic acid as a primary virus receptor.

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Introduction

Influenza virus infection is estimated to be responsible for approximately 36,000 deaths and 100,000 hospitalizations annually in the United States alone (Thompson et al., 2003, 2004). These numbers would increase considerably in the event of an influenza pandemic, which would result from the introduction of an influenza A virus strain encoding a hemagglutinin (HA) gene that was not present in viruses circulating recently in the human population (Barnett et al., 2005; Bartlett and Hayden, 2005; Wilson et al., 2005). Pandemic influenza A virus strains are generated through one of two mechanisms: reassortment or adaptation. Since the influenza A virus genome consists of eight distinct RNA segments (Lamb and Krug, 2001), infection of one cell with two influenza A virus strains can lead to the exchange of genetic material, a process called reassortment, resulting in a new virus strain possessing biological properties derived from either parental viruses (Horimoto and Kawaoka, 2005). The presence of influenza A virus strains bearing various combinations of the 16 HA and 9 neuraminidase (NA) serotypes in a range of animal reservoirs provides a pool of virus strains that could be the source of the next influenza pandemic (Fouchier et al., 2005; Rohm et al., 1996; WHO, 1980). The influenza pandemics of 1957 and 1968 resulted from reassortment of human and avian influenza A viruses (Webster et al., 1997). In contrast, the influenza pandemic of 1918 is believed to have been caused by an avian influenza A virus which acquired the ability to infect and efficiently spread in the human population through adaptation, not reassortment (Reid et al., 2004; Taubenberger et al., 2005; Tumpey et al., 2005).

The primary, but certainly not the only, hurdle influenza A virus must overcome when crossing a species barrier is believed
to be utilization of host cell receptors (Ito, 2000). Influenza A virus uses sialic acid (SA) residues present at the terminus of oligosaccharide modifications of host cell glycoproteins as attachment receptors (Chu and Whittaker, 2004). SA residues are usually linked to the penultimate carbohydrate group via α2,3-, α2,6-, or α2,8-linkages, and most influenza A virus strains characterized to date utilize either α2,3- or α2,6-SA as a primary receptor (Glaser et al., 2005; Matrosovich et al., 2000). Avian influenza A virus strains preferentially bind to α2,3-SA since this form of SA is abundant in the gastrointestinal tract of various avian species (Gambaryan et al., 2002, 2004; Kim et al., 2005; Matrosovich et al., 2000). However, while the human respiratory tract contains both α2,3- and α2,6-SA (Matrosovich et al., 2004; Shinya et al., 2006; Slepushkin et al., 2001; van Riel et al., 2006; Zhang et al., 2005), most human influenza A virus strains show a binding preference for α2,6-SA as a receptor (Baum and Paulson, 1990; Gagneux et al., 2003; Matrosovich et al., 2000, 2004).

In mammals, the primary tissues infected by influenza A virus are the trachea as well as the upper and lower bronchial tubes (Zambon, 2001). In rare cases, virus infection of the alveolar epithelial cells occurs, leading to viral pneumonia. Several animal models have been used to study influenza A virus pathogenesis, most notably the laboratory mouse (Lu et al., 1999; Novak et al., 1993). However, most human influenza A virus strains must be adapted for efficient replication in mice (Brown et al., 2001; Lu et al., 1999). Human influenza A virus clinical isolates replicate and cause disease in ferrets (Govorkova et al., 2005; Herlocher et al., 2001; Maher and DeStefano, 2004; Reuman et al., 1989; Sweet et al., 2002; Zitzow et al., 2002), Syrian golden hamsters (Ali et al., 1982; Daly et al., 2003; Murphy et al., 1997; Potter and Jennings, 1976; Renis, 1977; Snyder et al., 1989; Stein-Streilein and Guffee, 1986), cotton rats (Ottolini et al., 2005) and non-human primates (Rimmelzwaan et al., 2001), but with the possible exception of ferrets, none of these animal models is used extensively.

Since Syrian golden hamsters can be infected with influenza A virus and infected hamsters can transmit virus to uninfected animals, we characterized the ability of influenza A virus strains to infect primary, differentiated trachea epithelial cell (TEC) cultures derived from Syrian golden hamsters. Our results demonstrate that Syrian golden hamsters and hamster TECs represent an attractive system for studying infection of nonciliated airway epithelial cells, investigating the reassortment of influenza A virus strains which have dissimilar receptor preferences and generating influenza A virus strains with altered or expanded receptor utilization.

Results

Hamster TECs support influenza A virus replication

The ability of influenza A viruses to replicate in well-differentiated hamster TEC cultures grown in Transwell tissue culture inserts was determined by infecting the cultures with recombinant influenza A virus strains A/WSN/33 (rWSN; H1N1) and A/Udorn/72 (rUdorn;H3N2), and the clinical isolates A/California/7/2004 (H3N2), A/Memphis/14/96 (H1N1) and A/Memphis/5/98 (H3N2). Infectious virus titers in the supernatants of virus-infected cells were determined by plaque assay at various times post-infection. Hamster TEC cultures are routinely infected between days 10 and 14 after initiation of an air liquid interface (ALI), a time when the culture is fully differentiated into a polarized, heterogeneous cell population (Rowe et al., 2004). Both recombinant strains and clinical isolates displayed similar patterns of replication, consisting of a peak in viral titer at 1 to 2 days post-infection, and then a decline to undetectable amounts by 4 to 5 days post-infection (Figs. 1A, B). Virus was detected in the apical but not the basolateral supernatants over the time course of infection (Rowe and Pekosz, 2006), and proteases produced by the cells were responsible for cleavage of the HA protein into its active subunits (Rowe et al., 2004). Thus, laboratory strains and clinical isolates of influenza A virus replicate to equivalent infectious virus titers in hamster TECs.

The cell tropism associated with influenza A virus infection was determined by immunostaining virus-infected hamster TEC cultures for expression of the influenza A virus HA protein and
the ciliated cell marker β-tubulin IV after infection with rUdorn. Viral antigen expression correlated with the peak of virus titer, with antigen-positive cells being abundant at day 1 and decreasing in subsequent days post-infection (Fig. 2). Viral antigen appeared to predominate in the nonciliated cell population, particularly on day 1 post-infection, but limited numbers of viral antigen-positive, ciliated cells could be detected at day 2 post-infection. The decreased numbers of cell nuclei (visualized by TO-PRO-3 fluorescence) over time (Fig. 2, lower panels) suggested that virus infection induced cell death within the culture, most likely in the virus-infected cell population. However, virus infection did not result in death of all the cells in the culture, as demonstrated by the presence of ciliated cells at all times post-infection (Fig. 2, upper panels). These data demonstrate that the nonciliated cells in the hamster TEC cultures, are the primary cell type infected by rUdorn virus and that some subset of cells in the cultures are unable to support influenza A virus replication.

Expression of α2,3- and α2,6-linked SA in hamster TECs

Since influenza viral antigen expression was detected predominantly in the nonciliated cell population, we determined whether virus receptor expression could be responsible for the cell type-specific infection pattern. The expression of α2,3- and α2,6-linked SAs, the primary receptors for influenza A virus entry, was determined using lectins that recognize either α2,3- (Maackia amurensis agglutinin, MAA) or α2,6- (Sambucus nigra agglutinin, SNA) SA. Both α2,3- and α2,6-SA moieties were expressed (Fig. 3A), and quantitation of the number of cells per field expressing each linkage gave a ratio of 1.5:1 of α2,3- to α2,6-SA expressing cells (Fig. 3B). We next determined the distribution of α2,3- and α2,6-SA on specific cell types within the hamster TEC culture. α2,6-SA was localized to nonciliated cells, as demonstrated by the lack of colocalization of SNA with the ciliated cell marker protein β-tubulin IV (Fig. 3C, upper panels). α2,3-SA had a broader distribution than α2,6-SA, and was detected on both ciliated and nonciliated cells (Fig. 3C, lower panels). It is important to note, however, that the population of cells that expressed the highest levels of α2,3-SA were not β-tubulin IV positive (Fig. 3C, lower panels, white asterisks). Thus, the nonciliated cell population expressed both α2,3- and α2,6-linked SAs while the ciliated cells only express α2,3-SA but to a lower level than that observed on nonciliated cells.

rUdorn viral antigen was detected primarily in nonciliated cells at day 1 post-infection (Fig. 2), and this correlates with the high level of α2,3- and α2,6-linked SA expression in the same population of cells. To conduct a more thorough analysis of virus strain-specific differences in cell tropism, hamster TECs were infected with rWSN (H1N1) or rUdorn (H3N2), and viral antigen expression was analyzed during the first replication cycle (6 h post-infection), as well as 1 and 2 days post-infection. The rWSN virus has been passaged extensively in mice and embryonated eggs, conditions that have been shown to result in a preference for α2,3-SA as an entry receptor (Gambaryan et al., 1997, 1999; Matrosovich et al., 2000) and it encodes amino acids at positions 190 and 225 of the HA protein that dictate preferred binding to α2,3-SA (Glaser et al., 2005; Stevens et al., 2006). The rUdorn virus has been passaged in MDCK cells and MDCK cell passage is believed to maintain the α2,6-linked SA receptor preference of human influenza A virus strains (Gambaryan et al., 1997, 1999; Matrosovich et al., 2000). The amino acids present at positions 226 and 228 of the HA protein predict a preference for binding to α2,6 SA (Connor et al., 1994; Gambaryan et al., 1997, 1999, 2005; Govorkova et al., 1999; Ito et al., 1997; Rogers and Paulson, 1983; Rogers et al., 1983; Stevens et al., 2006); however, binding of rUdorn HA to α2,3 and α2,6-linked SA was detected using ganglioside binding assays (Ryan-Poirier et al., 1998).

At 6 h post-infection, rUdorn HA was expressed predominantly in the nonciliated cell population (Fig. 4A, a–c), and this same pattern is seen at day 1 post-infection (Fig. 4A, d–g). At day 2 post-infection, few viral antigen-positive cells are detected (Fig. 4A, i–k, white asterisks in k denote antigen-positive cells), suggesting that most of the virus-infected cells have succumbed to infection. Interestingly, almost all the antigen-positive cells detected at day 2 post-infection are ciliated. The numbers of viral antigen-positive cells were quantified on days 1 and 2 post-infection and data indicate that...
there are limited numbers (less than 5%) of infected, ciliated cells on both days post-infection (Fig. 4B). However, the majority of infected cells are not ciliated and the numbers of infected, nonciliated cells drop dramatically between days 1 and 2 post-infection. The numbers of ciliated cells in the infected cultures did not drop as significantly as the total number of nuclei, indicating that virus infection was eliminating nonciliated cells, consistent with the cell tropism data (Fig. 4C).

These data indicate that the primary cell type infected by rUdorn is the nonciliated cell, with some infected ciliated cells detected at days 1 and 2 post-infection. Since only α2,3-SA is detected in ciliated cells, the rUdorn virus is most likely utilizing this form of sialic acid to enter ciliated cells.

The Clara cell secretory protein (CCSP) was used as a marker for one type of nonciliated cell present in the hamster TEC cultures (Singh and Katyal, 2000). CCSP colocalized with rUdorn HA at 6 h post-infection (Fig. 4D, a–c), indicating that Clara cells are at least one type of nonciliated cell infected by influenza A virus. At day 1 post-infection, rUdorn HA expression was clearly detected (Fig. 4D, f), but the CCSP staining appeared slightly diminished, likely due to the cytopathic effects of virus replication on this cell type or to reduced translation of host cell proteins in virus-infected cells (Fig. 4D, e and g). Expression of α2,6-SA was evident at 6 h post-infection, but difficult to detect at day 1 post-infection (Fig. 4D, d and h) most likely due to the combined effect of death of the nonciliated cells and reduced SA expression due to the activity of the viral NA protein.

Viral antigen expression in rWSN-infected cells was similar to that of rUdorn at 6 h and day 1 post-infection, with HA detected predominantly in nonciliated cells (Fig. 4E, a–c and e–g). At day 2 post-infection, rWSN HA was detected in a limited number of ciliated cells; however, antigen-positive nonciliated cells were also present (Fig. 4E, i–k, ciliated cells marked with asterisks). The numbers of antigen-positive cells were quantified and the data again show that the nonciliated cells are the primary cell type infected by rWSN, with infected ciliated cells low in number (less than 5%) at days 1 and 2 post-infection (Fig. 4F). The reduction in number of nuclei per field was much greater than the reduction of ciliated cells per field, indicating that the nonciliated cells were preferentially eliminated in virus-infected monolayers (Fig. 4G). When the expression of α2,3-SA – the SA form expressed on ciliated cells – was assessed, the ciliated cells infected at day 2 showed reduced amounts of α2,3-SA when compared to nonciliated cells at 6 h or 1 day after infection (Fig. 4E, panels d, h, and l). These data suggest that rWSN infects primarily nonciliated cells, with limited numbers of ciliated cells infected at days 1 and 2 post-infection. Taken together, the data in Fig. 4 indicate that the expression of α2,3- and α2,6-SA on nonciliated cells correlates very closely with the predominant viral antigen expression at early and late times during infection with both rUdorn and rWSN viruses.

In addition to rWSN and rUdorn, clinical isolates A/California/7/2004, A/Memphis/14/96, and A/Memphis/5/98 were also examined for HA expression in the hamster TEC cultures at the same time points (Fig. 5). The A/Memphis/14/96 and A/Memphis/5/98 viruses have been documented to bind preferentially to α2,6-SA (Matrosovich et al., 2003, 2004). HA expression of all three clinical isolates displayed the same antigen pattern—expression of HA exclusively in the nonciliated cells during the first round of virus replication (Fig. 5A, a–c, e–g, i–k) and day 1, with little to no antigen detected at day 2. The number of antigen-positive cells, ciliated cells, and nuclei at day 1 and day 2 were quantified for each clinical
Fig. 4. Influenza A virus infection targets nonciliated cells in hamster TEC cultures. Cultures were infected with an MOI of 3, and were analyzed for antigen expression at the indicated times post-infection (A) rUdorn HA expression (red) compared to β−tubulin IV (green) (D) Colocalization of rUdorn HA (blue) with the secretory cell protein CCSP (red) and α2,6-SA (E) rWSN HA expression (red) compared to β−tubulin IV (blue) and α2,3-SA. White asterisks in panels A-k and E-k indicate virus-infected ciliated cells at day 2 post-infection. The same field is shown in each of the four panels for each row. Quantitation of antigen-positive cells (B, F) and ciliated cells and nuclei (C, G) at days 1 and 2 post-infection. Ten to fourteen fields were counted for each virus per day and the data are expressed as a percentage of the total cells (nuclei) present in each field. There are approximately 150–250 nuclei per field in uninfected cultures (Rowe et al., 2004). All images were taken at a 63× magnification.
Fig. 5. Influenza A virus clinical isolates exclusively target nonciliated cells in hamster TEC cultures. Cultures were infected with an MOI of 3 and were analyzed for antigen expression at indicated times post-infection. (A) Clinical isolates A/California/7/2004, A/Memphis/5/98, and A/Memphis/14/96 at 9 h post-infection. HA expression (red) compared to $\beta$-tubulin IV (green). The same field is shown in each of the four panels for each row. (B–G) Quantitation of antigen-positive cells (B, D, and F) and ciliated cells and nuclei (C, E, and G) at days 1 and 2 post-infection. Ten to fourteen fields were counted for each virus per day and the data are expressed as a percentage of the total cells (nuclei) present in each field. There are approximately 150–250 nuclei per field in uninfected cultures (Rowe et al., 2004). All images were taken at a 63× magnification.
isolate (Figs. 5B–G). While the number of ciliated cells stay relatively constant through 2 days post-infection, nuclei counts decrease, suggesting again that the nonciliated cell population is infected and eliminated after infection with influenza A virus (Figs. 5C, E, and G). However, in contrast to the laboratory strains, antigen-positive ciliated cells were not detected at any time point with the three clinical isolates (Figs. 5B, D, and F). Thus, it appears that the ability of rWSN and rUdorn to infect a limited number of ciliated cells is not shared by the clinical isolates of the influenza A virus strains analyzed, and may be related to altered SA receptor usage (Deom et al., 1986; Ohuchi et al., 1997; Ward, 1997), or that tissue culture passage has created minor populations able to bind to alternate receptors in the case of rUdorn. Clinical human isolates of influenza A virus, however, seem to maintain a more strict receptor specificity for the α2,6-SA expressing nonciliated cell population.

Discussion

Receptor utilization is an important determinant of viral pathogenesis and cell tropism. For influenza A virus, receptor utilization is particularly important as it is thought to be one of the key factors enabling avian influenza A viruses to replicate in and spread between humans (Parrish and Kawaoka, 2005). The ongoing outbreak of H5N1 virus occurring in poultry throughout Asia and parts of Europe has led to over 100 cases of human infection, thus increasing the likelihood that a variant explaining the poor ability of human influenza A virus strains to utilize different forms of SA (avian and human influenza viruses for example) as well as the adaptation of α2,3-SA utilizing viruses to α2,6-SA usage. The expression of α2,3-SA in ciliated cells was reduced when compared to expression in nonciliated cells, and ciliated cells were less susceptible to infection with rUdorn and rWSN. This may indicate that the absolute level of SA on the plasma membrane may also dictate the susceptibility of a cell to influenza A virus infection, but definitive statements on this await more careful and accurate quantification of SA levels in specific cell types. The effect of viral neuraminidase activity and specificity on the amount of available SA may also have a profound effect on cell susceptibility to virus infection.

The expanded cell tropism of rUdorn in hamster TEC cultures when compared to primary influenza A virus isolates may reflect the ability of the Udorn HA protein to recognize both α2,3- and α2,6-SA (Ryan-Poirier et al., 1998). Curiously, the Udorn HA protein possesses amino acid residues at positions 193, 226, and 228 that are found in H3 subtype viruses that bind preferentially to α2,6-SA (Medeiros et al., 2001, 2004; Stevens et al., 2006; van Riel et al., 2006). Position 186 of H3 subtype HA proteins has been implicated in altered receptor specificity with substitutions of G186V or S186I resulting in increased α2,3-SA binding (Gambaryan et al., 1999; Widjaja et al., 2006); however, the rUdorn strain encodes an S at this amino acid. This indicates that other, as yet unidentified amino acids may alter SA receptor recognition in H3 subtype HA proteins or that the assays used to detect α2,3- and α2,6-SA binding are not reflective of the cell tropism documented in this study. It is important to note that while receptor utilization and expression are important determinants of influenza A virus cell tropism in the respiratory tract (Matrosovich et al., 2004; Shinya et al., 2006; van Riel et al., 2006; Zhang et al., 2002, 2005), it has been demonstrated that interferon and interferon signaling are also important factors that limit influenza A virus tropism in vivo and in vitro (Garcia-Sastre et al., 1998a, 1998b).

The expression of α2,3- and α2,6-SA in the human respiratory tract has been investigated extensively (Matrosovich et al., 2004; Slepushkin et al., 2001; Zhang et al., 2005). Most studies support the model that α2,3-SA is present primarily, if not exclusively, in the ciliated cells. The expression of α2,6-SA has been documented to be either exclusively in the nonciliated cells (Matrosovich et al., 2004) or present in both ciliated and nonciliated cells (Ibricevic et al., in press; Zhang et al., 2005). The amount of SA on each cell type has yet to be quantified precisely. Highly pathogenic H5N1 influenza A virus strains have limited ability to attach to and replicate in the upper respiratory tract as compared to the lower respiratory tract—again the result (at least in part) of altered expression of α2,3- and α2,6-SA (Shinya et al., 2006; van Riel et al., 2006). The SA receptor pattern on hamster TEC cultures differs from that described for humans, an important factor to consider when interpreting hamster pathogenesis experiments. However, there are two points to consider when deciding between hamsters or mice for influenza pathogenesis studies. First, the hamster model may be better suited to pathogenesis experiments with human influenza A virus strains when compared to the mouse...
model since α2,6-SA is not expressed in the mouse respiratory tract, but is present in the upper and lower respiratory tract of the hamster (CMN and AP unpublished observations). Second, humans and hamsters have significant numbers of mucous-secreting goblet cells (Atherton et al., 2003; Rowe et al., 2004) while mice possess virtually undetectable numbers (Look et al., 2001; Shahzeidi et al., 2003; Walter et al., 2002). Since hamster and human goblet cells also express SA receptors, studies on the effects of influenza on this cell type are best performed in the hamster model.

It has been suggested that replication of influenza A virus in nonciliated cells is important for efficient virus replication and/or transmission in humans (Matrosovich et al., 2004). This may also explain the ability of hamsters to support the replication and transmission of human influenza A virus strains (Ali et al., 1982; Daly et al., 2003) and suggest that studying transmission of human and avian influenza A viruses in hamsters may help shed light on viral and host factors that are important for virus transmission. Our data demonstrate that primary, differentiated hamster TEC cultures can serve as a useful model of influenza A virus pathogenesis and cell tropism, and provide a relevant cell culture system for studying influenza virus strains that have different SA receptor specificities.

Materials and methods

Reagents and antibodies

The components in TEC basic media (TEC basic), proliferation media (TEC plus), and maintenance media (TEC MM) have been described previously (Rowe et al., 2004). Rabbit anti-hamster Clara cell secretory protein (CCSP; 1:500 immunofluorescence) was kindly provided by Gurmukh Singh (VA medical center, Pittsburgh, PA). Other primary antibodies were purchased and used as follows: mouse anti-β Tubulin IV (1:100 immunofluorescence; BioGenex, San Ramon, CA), goat anti-H3 hemagglutinin (HA-H3 subtype) Aichi/2/68 sera (1:500 immunofluorescence; NIH/NIAID reference reagent V314-591-157), goat anti-H1 hemagglutinin (HA-H1 subtype) A/PR8/34 sera (1:500 immunofluorescence; NIH/NIAID reference reagent V314-511-157). The influenza A virus HA antibodies showed minimal reactivity to mock-infected hamster TEC cultures (data not shown). Secondary antibodies were used as follows: donkey anti-mouse IgG conjugated to fluorescein isothiocyanate (1:250 immunofluorescence), donkey anti-goat IgG conjugated to rhodamine red (1:250 immunofluorescence), and goat anti-mouse IgG conjugated to rhodamine red (1:250 immunofluorescence) were purchased from Jackson Immunoresearch (Westgrove, PA); goat anti-mouse IgG conjugated to Alexa Fluor 647 (1:500 immunofluorescence), donkey anti-mouse IgG conjugated to Alexa Fluor 647 (1:500 immunofluorescence), goat anti-rabbit IgG conjugated to Alexa Fluor 594 (1:500 immunofluorescence), and TO-PRO-3 nuclear stain (1:500) were purchased from Molecular Probes (Eugene, OR); anti-digoxigenin-fluorescein, FAb fragment (1:50) was purchased from Roche applied science (Indianapolis, IN).

Cell lines

Madin–Darby canine kidney cells (MDCK American Type Culture Collection, Manassas, VA) were cultured in Dulbecco’s modified Eagle medium (DMEM, Sigma, St. Louis, MO) containing 10% fetal bovine serum (Atlanta Biologics, Atlanta, GA), 100 U/mL of penicillin, and 100 μg/mL of streptomycin (Invitrogen, Grand Island, NY) and maintained at 37 °C in a humidified environment containing 5% CO2.

Viruses

Recombinant influenza A viruses A/WSN/33 (rWSN) (Neumann et al., 1999; Takeda et al., 2002) and A/Udorn/72 (rUdorn) (Takeda et al., 2002), and the influenza A virus clinical isolates A/California/7/2004 (H3N2, Centers for Disease Control, Atlanta GA), A/Memphis/14/96 (H1N1), and A/Memphis/5/98 (H3N2) (St. Jude Children’s Hospital, Memphis, TN) were used in this study. The latter two virus strains have been characterized with respect to their cell tropism in human TEC cultures (Matrosovich et al., 2004) and receptor specificity (Matrosovich et al., 2003). Virus stocks were generated by infecting MDCK cells and infectious virus was quantified by plaque assay using MDCK cells as previously described (Paterson and Lamb, 1993). The H3 amino acid numbering system is used when referring to amino acid positions in HA.

Hamster TEC cultures

Hamster TEC isolation and culture conditions were performed as previously described (Rowe et al., 2004). In all experiments, hamster TEC cultures were seeded and maintained in 0.4 μM pore, 0.33 cm² Transwell–Clear support membrane (Corning Costar, Corning NY) (Rowe et al., 2004).

Infection of hamster TECs

Hamster TEC cultures were infected between days 10 and 14 after ALI. Hamster TECs were infected via the apical chamber with 1 × 10⁶ plaque forming units (pfu) of virus diluted in warm DMEM with penicillin/streptomycin in a total volume of 100 μL. If all the cells in the culture were susceptible to influenza virus infection, this would correspond to a multiplicity of infection (MOI) of approximately 3. Since the cultures are pseudostratified, not all of the cells in the culture will be exposed to the apically administered virus inoculum. Also, not all cell types express influenza A virus receptors. Cells were incubated with virus at 37 °C for 1 h, inoculum removed, and cells washed three times with 200 μL of warm DMEM with penicillin/streptomycin. After washing, 100 μL of DMEM with penicillin/streptomycin and 500 μL of TEC maintenance media (MM) (Rowe et al., 2004) were placed in the apical and basolateral chambers, respectively. Apical and basolateral supernatants were collected at the indicated times post-infection and stored at −70 °C. All growth curves and immunostaining on hamster TEC cultures were reproduced at least three times.
**Immunofluorescence confocal microscopy**

At indicated times post-infection, hamster TECs were washed three times with phosphate buffered saline (PBS, Gibco Inc., Carlsbad, CA), and fixed in PBS containing 2% paraformaldehyde for 15 min at room temperature. Cells were washed three times and permeabilized with PBS containing 0.2% Triton-X 100 and 0.1% sodium citrate for 10 min at room temperature. Cells were washed with PBS and incubated in PBS containing 3% normal goat or normal donkey serum and 0.5% bovine serum albumin (blocking buffer) for 30 min at room temperature. Cells were washed and incubated with blocking buffer containing primary antibodies for 1 h at room temperature, washed again, and incubated with blocking buffer containing secondary antibodies and TO-PRO-3 for 45 min. The wash solution for all steps is PBS with 0.2% Tween-20.

**Detection of lectin binding**

The DIG glycan differentiation kit (Roche, Indianapolis, IN) was used. SA linked to the penultimate galactose or N-acetylgalactosamine via an α2,6 linkage was detected with digoxigenin-conjugated *M. amurensis agglutinin* (MAA) lectin while α2,6 linked SA was detected with digoxigenin-conjugated *S. nigra agglutinin* (SNA) lectin. Samples were stained according to the manufacturer’s protocol using anti-digoxigenin FITC. Lectin-positive cells in 10 to 12 fields were counted to determine the number of cells expressing each linkage. There are between 150–250 total cells per field, as determined by TO-PRO-3 staining. Lectin staining on hamster TEC cultures was reproduced in three separate experiments.

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