Entry of parainfluenza virus into cells as a target for interrupting childhood respiratory disease

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The paramyxovirus family of viruses and the parainfluenza viruses

Viruses belonging to the paramyxovirus family, particularly respiratory syncytial virus (RSV), the recently identified human metapneumovirus (1), and the human parainfluenza viruses (HPIVs) types 1, 2, and 3, cause the majority of childhood cases of croup, bronchiolitis, and pneumonia worldwide (2). HPIV3 alone is responsible for approximately 11% of pediatric respiratory hospitalizations in the US (3, 4) and is the predominant cause of croup in young infants, while HPIV1 and -2 tend to infect older children and adolescents. While other causes of respiratory disease in children—influenza and measles—have yielded in part to vaccination programs and antiviral therapy, children are still virtually unaided in their battle against the major causes of croup and bronchiolitis. RSV has been extensively studied, and some effective strategies of prophylaxis have been developed (5), but for the parainfluenza viruses, there are no therapeutic weapons; advances in preventing and treating diseases caused by both groups of viruses, especially the parainfluenza viruses, are far behind those in combating diseases caused by many more genetically complex pathogens.

The parainfluenza viruses replicate in the epithelium of the upper respiratory tract and spread from there to the lower respiratory tract. Epithelial cells of the small airways become infected, and this is followed by the appearance of inflammatory infiltrates. The relationship among the tissue damage caused by the virus, the immune responses that help to clear the virus, and the inflammatory responses that contribute to disease is still quite enigmatic. Both humoral and cellular components of the immune system appear to contribute to both protection and pathogenesis (6, 7). Infection with HPIV in immunocompromised children (e.g., transplant recipients) is associated with a range of disease, from mild upper-respiratory symptoms to severe disease requiring mechanical ventilation and leading to death (8).

The hurdle for developing modes of preventing and treating croup and bronchiolitis caused by parainfluenza has been in large part a result of the gaps in our understanding of fundamental processes of viral biology and of the interaction of these viruses with their hosts during pathogenesis. For example, an inactivated HPIV1, -2, -3 vaccine used in infants in the late 1960s was immunogenic but did not offer protection from infection (9, 10), which highlights the challenge of identifying which elements of the immune response confer protection from HPIVs. Primary infection with any HPIV does not confer permanent immunity against that virus, and repeated reinfection with the same agent within a year of the previous infection is common in young children. Immunity generated after the first infection is, however, often sufficient to restrict virus replication in the lower respiratory tract and prevent severe disease. Efforts are currently underway to develop live attenuated vaccines against HPIV1, -2, and -3, and an increased understanding of the molecular basis for attenuation of virulence may eventually lead to live HPIV vaccines that can be designed to be both attenuated and immunogenic and even to the development of combination respiratory virus vaccines (reviewed in ref. 11). Deeper understanding of the interplay among virus-mediated pathology, beneficial immune responses, and exaggerated or disease-enhancing inflammatory responses will be vital for developing safe and effective vaccine strategies.

Antiviral therapy for the parainfluenza viruses has not been explored but, in light of the complexities involved in vaccination, could be a principal weapon against these diseases. Several features of the viral life cycle make these viruses vulnerable to attack. HPIVs enter their target cell by binding to a receptor molecule and then fusing their viral envelope with the cell membrane to gain admittance to the cytoplasm. Binding, fusion, and entry are therefore critical stages at which we could interfere with the viral life cycle and prevent disease. A firm grasp of the molecular mechanisms of these events is the basis for understanding respiratory disease pathogenesis and developing potential approaches to prevention and treatment.

The parainfluenza virus life cycle

HPIVs are members of the Respirovirus and Rubulavirus genera within the Paramyxoviridae family. The viruses are roughly spher-
Virions are formed, according to the prevailing model for virion assembly, when newly assembled nucleocapsids containing the full-length viral RNA genome along with the polymerase proteins bud out through areas of the plasma membrane that contain the F and HN proteins and the matrix protein. In polarized epithelial cells, the viruses bud from the apical surface of the cell. The matrix protein binds to the nucleocapsid and also interacts with the cytoplasmic tails of the HN and F proteins, in this way mediating the alignment of the nucleocapsid with the areas of the plasma membrane containing viral glycoproteins in order to set the scenario for budding (16). The neuraminidase or receptor-cleaving activity of the HN molecule cleaves sialic acid–containing receptor moieties that would attach the viral HN protein to the cell surface and allows the release of newly budded particles from the cell to begin a new round of infection (17, 18).

**Role of the parainfluenza surface protein HN in receptor binding, receptor cleaving, and F protein activation to mediate fusion**

The HN proteins of HPIVs are different from the receptor-binding glycoproteins of other members of the paramyxovirus family in that they possess both hemagglutinating (sialic acid–containing receptor–binding) and neuraminidase (sialic acid–containing receptor–cleaving) activities. The parainfluenza HN proteins are oriented such that their amino termini extend into the cytoplasm, while the C termini are extracellular (Figure 1). The HN protein is present on the cell surface and on the virion as a tetramer composed of disulfide-linked dimers (19). The molecule contains a cytoplasmic domain, a membrane-spanning region, a stalk region, and a globular head. Crystal structures of the HN protein of the avian paramyxovirus known as Newcastle disease virus (NDV) (20, 21) and more recently the HPIV3 HN protein (22) demonstrate that the globular head contains the primary sialic acid–binding site and the neuraminidase active site.

Far from simply serving to attach the virus to the surface of the cell and to release virus after replication, the interaction of the HN protein with its receptor is required for F protein–mediated membrane fusion during viral infection (23, 24). Studies of several related paramyxoviruses have revealed that, for most members of this family, the HN protein is essential to the F protein–mediated fusion process (25–27). While receptor binding is an important component of this process, attachment is not sufficient (23, 25, 28); many F proteins demonstrate a requirement for the presence of an HN protein from the same type of virus (the homotypic HN protein) in order to mediate fusion (25, 26). One proposed explanation for this requirement is that the interaction between the HN and F proteins may be type specific and/or that a specific relationship between the structures and/or activities of the 2 proteins is required in order to maintain function (29). This final key function of the HN protein — promotion of fusion — has become amenable to mechanistic study only recently; upon binding to its receptor, parainfluenza HN protein plays a critical role in activating or “triggering” the F protein to assume its fusion-ready conformation (30, 31). Since insertion of the fusion peptide region of the F protein into the target cell membrane after the activation step is the key event leading to membrane fusion, efficiency of F protein triggering by the HN protein is an important variable influencing the extent of fusion mediated by the F protein and thus the extent of viral entry.

**Triggering of fusion during entry by enveloped viruses**

Entry of all enveloped viruses into host cells requires fusion of the viral and cell membranes. The fusion protein that mediates these processes differs among the enveloped viruses, but thus far these have been mechanistically grouped into just 2 classes of proteins.
The first, termed class I (reviewed in ref. 32), includes the paramyxovirus fusion proteins as well as the influenza hemagglutinin protein, the HIV gp120 fusion protein, and the Ebola virus fusion protein. Each is synthesized as a single polypeptide chain that forms trimers and is then cleaved by host proteases into 2 subunits, exposing a fusion peptide that will insert into the target cell membrane (33, 34) (reviewed in ref. 35). The trigger that initiates a series of conformational changes in the F protein leading to membrane merger differs depending on the pathway the virus uses to enter the cell and thus whether fusion needs to occur at the surface at neutral pH or in the endosome. The influenza HA protein has been the most extensively studied model for class I fusion (36), and the conformational change is triggered by the acidic pH of the endosome, which then allows the viral and endosomal membranes to fuse (35). Class II fusion proteins include the flavivirus dengue virus E protein (37), tick-borne encephalitis virus E protein (38), and togavirus Semliki Forest virus E1 protein (39), and despite pronounced differences in the structures of class I and class II fusion proteins, their transition to the post-fusion state proceeds through structures similar enough to suggest a common mechanism (40).

The paramyxovirus fusion process is thought to occur at the surface of the target cell at neutral pH, and activation of the F proteins occurs when the adjacent HN protein binds to the sialic acid-containing receptor, permitting fusion to occur. For HPIV3, the binding of HN protein to its receptor triggers the F protein to fuse with the target cell membrane, and alterations in the HN protein can alter its ability to trigger the F protein (30). The fusion peptides, which are buried within the F protein trimer, must be exposed in order to insert into the target cell membrane, and additional coreceptor binding events, for either the HN or F protein, have not been ruled out. New structural and experimental information about paramyxovirus F proteins has led to models for the structural transitions that occur during class I fusion (31, 32, 40–42) (Figure 3). The ectodomain of the membrane-anchored subunit of the F protein contains 2 hydrophobic domains, the fusion peptide and the transmembrane-spanning domain. Each of these domains is

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**Figure 2**
A schematic illustration of the parainfluenza viral life cycle. RER, rough endoplasmic reticulum.
adjacent to 1 of 2 conserved heptad repeat (HR) regions: the fusion peptide is adjacent to the N-terminal HR (HR-N), and the transmembrane domain is adjacent to the C-terminal HR (HR-C). These HR domains can oligomerize into coiled coils composed of several \( \alpha \)-helices. Once the F protein is activated, the fusion peptide inserts into the target membrane, first generating a transient “prehairpin” intermediate that is anchored to both viral and cell membranes (Figure 3, A–C). This form then refolds and assembles into a fusogenic 6-helix bundle structure as the HR-N and HR-C associate into a tight complex with N- and C-peptides aligned in an antiparallel arrangement (Figure 3, D and E). The resultant helical coiled-coiled rods are located adjacent to the fusion peptides, forming a highly stable 6-helix bundle (Figure 3F). Refolding relocates the fusion peptides and transmembrane anchors to the same end of the coiled coil, bringing the viral and cell membranes together. The formation of a coiled-coil structure during this step generates the free energy for the membranes to bend toward each other and is thought to be the driving force for membrane fusion (31).

The role of the HN protein in activating the F protein has been explored using peptides that correspond to the HR domains of the F protein. Synthetic peptides derived from the HR regions of several paramyxovirus F proteins can inhibit fusion by binding to their complementary HR region and thereby preventing HR-N and HR-C from refolding into the stable 6-helix bundle structure required for fusion (32, 41). Susceptibility to inhibition by these peptides can be used as a gauge of F protein’s progress through the steps outlined above, and such studies suggest that binding of the HN protein to a sialic acid–containing receptor induces a conformational change in the F protein (31). One model for the triggering of fusion suggests that, upon receptor binding, the HN protein itself undergoes a receptor-induced conformational change, which in turn triggers the conformational change in the F protein (14, 24). Results of experiments using F proteins that fuse without the HN protein suggest that the presence of the HN protein lowers the energy of activation required for F protein–mediated fusion (43). Many of the fundamental aspects of this fusion activation process remain to be understood; in particular, why is there a requirement for HN–receptor binding in order to initiate fusion promotion? Does the HN protein undergo conformational change upon receptor binding? An understanding of how the HN protein carries out its F protein–triggering function is central to understanding paramyxovirus entry.

Use of variant HN proteins to scrutinize the HN-triggered F protein–mediated fusion process

We have focused on the process whereby HPIV3 HN protein triggers the F protein after receptor interaction in order to understand this step in the entry of HPIV3 into the host cell. Since the HN protein has 3 functions (receptor binding, neuraminidase, and F protein activation) that each impact the fusion potential of the virus, the field had been hindered by the inability to study just 1 function — triggering — independently of the influence of other HN protein activities. A strategy for quantitating F protein triggering was developed using a panel of mutant HN proteins (17, 18, 24, 30) in order to map the 3 functions of the HN protein to specific regions of the protein. One of the mutant HN proteins displayed a defect in triggering the F protein, which demonstrated for the first time that F protein triggering is a distinct function of the HN protein (30). In our assays, insertion of F protein into target rbc membranes — or fusion between cells — served as a surrogate for the first steps of viral entry into target cells; while these systems differ from the setting of natural infection in the human lung, they allow representation of the events of HN-receptor binding, F protein triggering, and F protein insertion, and mechanistic models can then be tested in either epithelial cell cultures or animal models.

Viruses with alterations in the HN protein that led to either an increased affinity of the protein for its receptor or decreased neuraminidase activity are more potent at inducing membrane fusion, which leads to the conclusion that it is the interaction of the HN protein with its receptor that is required for activation of F protein (29). HN proteins with either increased receptor affinity or decreased enzymatic (receptor-cleaving) activity stay engaged with their receptor longer than WT HN proteins (29). Both cell surface receptor–binding and neuraminidase activities of the HN protein, by impacting the duration of HN-receptor engagement, regulate F protein activation and fusion promotion. An intriguing mutant virus is one containing an alteration (P111S) in the stalk region of the HN molecule along with a globular head mutation that decreases neuraminidase activity. While viruses with this HN molecule are indeed neuraminidase-deficient, they are defective in promoting cell fusion, in contrast to what might be predicted (30). This was the first indication that there is a separate HN protein function that specifically triggers F protein and that it resides in the stalk region of the protein.

The assay for measuring the F protein–triggering function of the HN protein (30) takes advantage of a sialic acid–containing receptor analog/neuraminidase inhibitor compound, 4-guanidino-Neu5Ac2en (4-GU-DANA; zanamivir, Relenza) that serves as a clinically effective antiinfluenza agent (44). While 4-GU-DANA inhibits HPIV neuraminidase activity, it does not prevent release of virus from the infected cell surface (45) as it does in the case of influenza viruses; instead, it blocks interaction between the parainfluenza HN protein and its receptor and thus — surprisingly — aids in the release of newly assembled virions from the infected cell (18). However, by interfering with HN-receptor interaction, 4-GU-DANA blocks receptor binding and thereby blocks fusion and viral entry (18, 45). These findings have stimulated interest in designing binding/entry inhibitors for treatment of paramyxovirus infection.

When WT and variant HN proteins were compared for their ability to trigger WT F protein, several findings emerged. The HN protein derived from the neuraminidase-deficient/fusion-defective virus has 2 amino acid alterations (D216N, in the globular head, and P111S, in the stalk) and is slower at activating F protein than either the WT HN protein or the singly mutated D216N HN protein. Comparison with a singly mutated P111S HN protein revealed that this triggering delay is entirely attributable to the P111S mutation (30). F protein triggering was dramatically reduced by this change in the stalk region of the molecule, although there was no decrease in receptor binding avidity. Conducting the experiments at a temperature and pH not permissive of neuraminidase activity eliminated the effects of the HN protein’s neuraminidase activity. As a result of the diminished triggering, cell fusion was also markedly reduced. The virus containing the doubly mutated D216N/P111S HN protein is the only paramyxovirus variant virus found to be specifically defective in the HN protein’s F protein–triggering and fusion promotion function, and this was the first time that a fusion defect could be specifically attributed to the HN protein’s triggering function, independent of the effects of the other 2 HN protein functions (30). The fact that 1 change in the HN protein led to a specific defect in fusion promotion showed that it is, indeed, the HN protein that activates the F protein.
Properties of the HN protein that modulate its ability to trigger the F protein

When known variant HN proteins are studied under experimental conditions that allow assessment of all 3 HN protein functions, it becomes evident that the balance among these properties determines entry (29). F protein triggering by the WT HN protein is dramatically reduced at a pH close to the optimum for neuraminidase (pH 5.7); target cell receptors are released from HN protein by neuraminidase, and little triggering occurs. For the neuraminidase-dead HN protein (D216N/P111S HN), however, the rate and extent of F protein triggering are the same at both pH 5.7 and pH 8.0 (a pH at which the neuraminidase is not active), which confirms that for the WT HN protein, it is the enhanced neuraminidase activity at low pH that diminishes F protein triggering. Comparison of the doubly mutated protein with the P111S HN (30) revealed the effect of the HN protein’s neuraminidase activity: the P111S HN (with residual neuraminidase activity) releases reversibly bound receptors from the HN protein, and unlike the D216N/P111S HN, its triggering cannot “catch up” by remaining in longer contact with receptor. A pH conducive to increased neuraminidase activity (pH 5.7) completely abolishes triggering by the P111S HN. This comparison of different HN proteins — with neuraminidase activity as the only variable — illustrates the key role of this enzyme in regulating F protein triggering: neuraminidase reduces the chance that the HN protein will remain in contact with target cell receptors and thus prevents the slowly triggering HN protein from performing.

To assess the impact of receptor avidity on triggering, a variant HN (T193A) with higher avidity for receptor than the WT HN, and with WT neuraminidase activity, was useful. For this variant, F protein triggering remained as high at pH 5.7 as at pH 8.0, and target cell receptor release remained as low at pH 5.7 as at pH 8.0, which suggests that higher receptor avidity counterbalances the effect of increased neuraminidase. Both neuraminidase activity and receptor-binding avidity impact receptor availability and thereby the efficiency of the third function, F protein triggering. Thus, while mutations in the stalk region (e.g., P111S) affect the triggering potential of the HN protein, expression of this potential is also modulated by alterations in the globular head that affect HN-receptor interaction. Triggering absolutely depends on HN-receptor interaction, and each of the 3 discrete properties of the HN protein independently affect the ability of the HN protein to complement the F protein in mediating fusion.
Probing the active sites on the HN protein

Since HN-receptor interaction influenced by receptor avidity and neuraminidase activity in the globular head determines the possible extent of triggering, this domain would be a prime target for antiviral drugs and key to HN protein function. 4-GU-DANA blocks the receptor-binding and neuraminidase functions of the HN protein (44), and therefore variant HN proteins that are resistant to this inhibitor reveal information about the site or sites on the HN protein responsible for these functions.

Resistance to inhibitors at the binding site. Just when experimental data had allowed us to generate predictions about the mechanism of 4-GU-DANA resistance in HPIV3 as well as potential differences between HPIV3 and NDV, Lawrence et al. obtained the crystal structure of the HPIV3 HN protein (22). The structure of the globular head region is shown in Figure 4A complexed with sialic acid. A single alteration in the HN protein – T193I – leads to an HPIV3 variant with phenotypic resistance to the effects of 4-GUDANA in terms of both neuraminidase activity and receptor binding (45). Increased receptor-binding avidity alone can confer drug resistance and indeed accounts for part of the variant virus’s 4-GU-DANA-resistant properties (45). However, the T193I substitution does not confer resistance to a smaller molecule, DANA (identical to 4-GU-DANA except for a smaller substitucent group at C4). In addition, substitution of a (smaller) alanine for the threonine in the active site (to generate T193A HN) does not confer resistance to 4-GU-DANA. It thus seemed possible that substitution of the larger isoleucine for threonine in the active site (at residue 193) might be excluding the inhibitor molecule from the active site of the resistant variant and contributing to resistance. Analysis of the crystal structure indeed shows that a T193I alteration in the HPIV3 HN protein would likely place the side chain of the isoleucine in a conformation that could lead to steric clash between the isoleucine at 193 and the guanidinium moiety of the 4-GU-DANA. Figure 4B shows the active site region complexed with 4-GU-DANA and reveals the extension of the guanidine moiety into the pocket. The structure thus supports the notion that part of the resistance of the T193I HN variant to 4-GU-DANA indeed arises from a reduction in binding of 4-GU-DANA due to the bulk of the isoleucine side chain. This pathway whereby HPIV3 could develop resistance to such compounds is an issue that needs to be carefully considered in the design of antiviral analogs.

An alteration at the HN protein dimer interface that affects avidity. One HPIV3 variant HN protein (H552Q HN) is resistant to 4-GU-DANA solely due to its higher avidity for the receptor; this HN protein is like the WT HN protein in its neuraminidase activity and neuraminidase sensitivity to 4-GU-DANA. H552 lies at the HN dimeric interface (22) and does not appear to be involved in forming the primary receptor-binding site, which is consistent with the fact that the mutation has no effect on 4-GU-DANA binding affinity or on neuraminidase activity. How can the increased receptor-binding avidity of the H552Q variant (24, 45) be explained? Either the mutation causes an indirect conformational change at the binding site or H552 could represent part of a second receptor-binding site. The possible existence of a second binding site for the HPIV3 HN protein at or near the dimer interface is under study.

HN molecules from related paramyxoviruses differ in their response to receptor analog inhibitors. The 3D crystal structure of the HN protein of the avian paramyxovirus NDV (20) suggests that one site could carry out both binding and neuraminidase activities, but researchers have postulated that an additional HN receptor-binding site exists (21), one exhibiting enzyme and receptor-binding activity and the second exhibiting only receptor binding activity (46). In NDV, as in HPIV3, 4-GU-DANA drastically reduces infection. However, in the case of HPIV3, this is an effect mediated by the blocking of viral entry into the host cell, whereas in the case of NDV, the binding of the HN protein to its receptor is resistant to the inhibitory effects of 4-GU-DANA, which suggests that failure of progeny virion release due to neuraminidase inhibition by 4-GU-DANA accounts for the reduced infectivity for NDV (46). Thus, even 2 similar paramyxoviruses behave entirely differently in terms of their response to receptor analog inhibitors, a finding central to the discussion of antiviral approaches (discussed below).

The availability of the HPIV3 HN protein crystal structure and data from our mutant HN protein studies have allowed several correlations between HPIV3 HN structure and function to be made. Residue T193 forms part of the primary active site in the globular head of the HN protein, and alterations at this site can affect both receptor and inhibitor binding. H552 modulates avidity of the HN protein for its receptor but does not form part of the primary receptor-binding site, lying instead at the dimer interface. Residue D216 forms part of the framework of the globular head active site region, in line with the observation that it is key for neuraminidase activity (Figure 4A). Mutations at any of these sites may have the potential to alter fusion promotion, via alteration of the period of time that the HN protein and its receptor are in contact (which is essential for F protein activation), which emphasizes the relationship among the 3 HN protein properties that contribute to entry.

Contribution of HN-receptor interaction to pathogenesis in vivo

For HPIVs, the interplay among virus-mediated pathology, beneficial immune responses, and disease-enhancing inflammatory responses is not well understood, and it is likely that, as for RSV (47), in many cases, disease severity is increased and the pathology of clinical disease is actually caused by the inflammatory response rather than by the cytopathic effects of the virus (47). This fundamental concept is highlighted by the fact that virus titers in the infected host are generally waning by the time disease symptoms become apparent (2) and that virus titer does not correlate with the severity of lower-respiratory disease. A cotton rat model of disease has proven useful in initial analyses of the factors affecting the pathogenesis of HPIV3 in vivo. Experimental infection of the cotton rat leads to infection of bronchiolar epithelial cells and to bronchiolitis, mimicking human disease, which makes this a relevant model for HPIV3 lower-respiratory infection (48). In a study of cotton rats infected with WT HPIV3 and 3 of the variant HN viruses described above — HN T193A (high receptor avidity, globular head mutation), HN H552Q (high receptor avidity, dimer interface mutation), and HN D216N (low neuraminidase activity, globular head mutation) (49) — there was normal clearance of the variant viruses compared with WT viruses, the variant plaque morphology was preserved in vivo, and there was no reversion to WT phenotype in the infected animals. Quite surprisingly, each of the HN protein alterations led to striking differences in the ability of HPIV3 to cause extensive disease, and this effect was dissociated from effects on viral replication. The variants caused alveolitis and an interstitial infiltrate, while the WT virus only caused peri-bronchiolitis. The enhanced disease caused by the HN variants was manifested by greatly increased inflammatory cell infiltrate in the alveoli and interstitial spaces in the lung, characterized by notably
thickened alveolar walls and marked recruitment of inflammatory cells within the air spaces. These results suggest that these differences are indeed due to modulation of the inflammatory response through the different HN protein activity of the variants and are dissociated from viral replication or infectivity.

We hypothesize that changes in the HN protein that alter either the affinity of the HN protein for receptor or receptor-cleaving activity may alter the nature of the inflammatory response of the host. Using HN variants to dissect the etiology of enhanced disease, it may be possible to identify which component(s) of the immune system's response to HPIV3 contributes to disease. Indeed, preliminary experiments suggest that the enhanced pathology observed following infection of the cotton rat with HN-variant HPIVs correlates with specific alterations in the chemokine response to infection that are distinct for each variant HN protein (50). If further experiments support the finding that HN protein alterations specifically alter chemokine expression, this will provide information about the immune contribution to pathogenesis that can be used to develop therapies to modulate an overactive inflammatory response following HPIV3 infection.

For influenza, the severity of disease may be related to the ability of individual strains to induce proinflammatory cytokine expression (51, 52), and cytokine levels appear to correlate with severity of illness (53–55). In a mouse model of disease, it is the HA protein (receptor-binding protein) of the highly virulent 1918 influenza “Spanish flu” that confers the ability to cause severe disease; the disease (as in the cotton rat experiments described above) was widespread and involved recruitment of neutrophils to the alveoli, while viruses with WT HA protein led to only limited involvement of the alveoli, an effect not attributable to differences in replicative ability (56). These findings correlated with greatly enhanced cytokine production, which suggests that this specific HA protein is a critical determinant of macrophage activation and of production of neutrophil chemoattractants. These findings are reminiscent of the enhanced disease caused by the HPIV3 HN variants.

**Strategies for blocking fusion and viral entry**

Drawing on all that is known about entry into the cell by HPIV and other enveloped viruses, a number of potential strategies for influencing viral fusion become evident: first, blocking or perturbing F-triggering, and second, blocking HN protein–receptor binding. Both events would result in failure of the virus to enter the target cell. As mentioned earlier, peptides derived from the HR-N and HR-C regions of class I F proteins (called HR-N and HR-C peptides) can interfere with fusion intermediates of the F protein (41, 57–61). For example, the HIV envelope glycoprotein gp160 attaches to cellular receptors via its gp120 subunit and mediates fusion via its gp41 subunit; HIV peptides corresponding to the HR-C domain of gp41 are effective for treatment of HIV in humans, and T-20 was the first synthetic HR-C peptide approved for HIV treatment (62, 63). The C-terminus of the HR-N trimer contains a hydrophobic pocket that provides a potential binding site for small molecules that might disrupt the hairpin structure (64) and could provide advantages over peptides for clinical use. Intriguingly, a low-molecular-weight molecule that is highly effective in inhibiting RSV fusion was recently shown to bind within this hydrophobic pocket of HR-N, which suggests that indeed a small molecule that disrupts the hairpin can derail the fusion process (65); similar results have been obtained for the paramyxovirus simian virus 5, which suggests the general applicability of this approach (66). Inhibition of the F protein triggering process, by peptides or other small molecules that interact with the HR regions, is a promising area for development of antiviral therapies.

While F protein activation is key for entry, the correct timing of F protein activation is also essential; triggering must occur when the F protein is in contact with the target cell membrane. A fusion inhibitor effective against influenza was shown to prematurely trigger the conformational change in HA protein, rendering the virus incapable of fusion (67). A similar mechanism has been proposed for HIV (32, 68). We suggest (29) that correct timing of activation, which for HPIV3 must depend on the balance among the HN protein’s receptor-binding, receptor-cleaving, and triggering
activities, is critical to entry and represents a potential target for intervention. Swaying the balance of HN protein activities toward premature triggering of parainfluenza F protein may be a strategy for preventing entry.

Since HN-receptor interaction is the critical prelude to F protein triggering, it is an attractive step for blocking viral entry, and the recent availability of 3D structures for NDV and HPIV3 would seem to make it more feasible to design inhibitors that specifically fit into the binding pocket on the globular head of the HN protein. For HPIV3, sialic acid analogs such as 4-GU-DANA, while they do inhibit neuraminidase (18), counteract infection by inhibiting receptor binding. It is thus possible that for HPIV3, sialic acid analogs may be viable antiviral agents by function as binding/entry inhibitors. For NDV, however, recent results (see above) indicate the opposite (46); these compounds inhibit neuraminidase and reduce infection in culture but do not completely prevent binding or block viral entry. It is possible that sialic acid analogs that are specifically designed to inhibit the active site of NDV neuraminidase may inhibit virion release as they do in the case of influenza virus (69). These data encourage optimism that receptor blockade seems to make it more feasible to design inhibitors that specifically target virus has been isolated after treatment of immunocompetent children. This number rose to at least 4% for treated adults harboring influenza viruses with oseltamivir-resistant neuraminidases, this number rose to at least 4% for treated children. Recently, a small study of children in Japan (88) found that, out of 50 children treated with oseltamivir, 9 (18%) harbored viruses with drug-resistance mutations in the neuraminidase gene; the mutations were located where predicted from the in vitro studies discussed above. These mutations occurred far more frequently than has been previously observed (89), but it has not yet been established whether this is a general phenomenon, nor whether the oseltamivir-resistant viruses are transmissible or pathogenic (90).

This tale, both in the elements of resounding success and in the elements of increasing concern about the development of resistance (88), points to the great utility of structural and in vitro studies brought to bear on development of antiviral therapies. In order to continue to benefit from these potent antiviral compounds, it is critical to understand more about which features of neuraminidase inhibitors will discourage the emergence of resistance (90).

Inhibition of neuraminidase: how HPIV is different from influenza and what can be learned from the influenza experience

While neuraminidase inhibition does not seem a promising strategy for interfering with HPIV infection, this strategy has met with success in treating influenza virus infections. For influenza, HA protein, which recognizes the sialic acid moiety on the cell surface receptor, mediates both receptor binding of the virus to the cell and fusion of the viral envelope with the endosomal membrane; the neuraminidase protein (NA) is necessary for promoting the release of newly formed virions from the cell surface because it removes receptors for the virus, preventing self-aggregation (70). While in the case of HPIV infection, 4-GU-DANA interferes with HN-receptor interaction and thus actually enhances virus release (18), in the case of influenza virus infection, the clinical effectiveness of this molecule has been attributed to its ability to halt spread of the virus when given early in infection (71, 72). We have found that sialic acid–based inhibitors of influenza virus NA can also exert a direct effect on the function of the other envelope protein, HA protein (44). Recent experiments in primary cultures of human airway epithelium cells demonstrated that oseltamivir (a sialic acid analog related to 4-GU-DANA, discussed below) interfered with influenza infection at the early stage of entry (73). Thus, while the effects of 4-GU-DANA on influenza virus have been ascribed purely to the prevention of viral release by neuraminidase inhibition, these results suggested that the antiviral mechanism of action of 4-GU-DANA might be broader and may extend to interfering with viral entry (44). It will be of great interest to determine whether, as is possible for HPIV, neuraminidase may play a role in early infection and whether inhibition at this stage of the viral life cycle contributes to clinical effectiveness.

The design of 4-GU-DANA as a sialic acid analog antiviral compound that mimics the virus’s natural substrate proceeded directly from 3D structural studies of the influenza NA (69). Zanamivir is administered by oral inhalation, which delivers the drug directly to the respiratory tract epithelium, and is clinically effective if given early in infection, with remarkably few side effects (74). Shortly after the introduction of zanamivir in clinical practice, an orally available NA inhibitor, oseltamivir, was developed. The NA inhibitors as a class are effective against all NA subtypes and therefore against all strains of influenza (74–82), including the 2004 avian influenza H5N1 strains that are resistant to the M2 inhibitors (83, 84).

An important aspect of the utility of these compounds is that until recently, there seemed to be very little development of resistance to neuraminidase inhibitors (85). The structure-based design of the neuraminidase inhibitors contributed to the fact that it is unlikely for the viral neuraminidase to change in such a way as to confer resistance, while still maintaining function. The neuraminidase inhibitors must fit directly into the enzyme’s active site pocket in order to block the enzyme’s activity, and since zanamivir was designed to closely resemble the natural substrate, mutations that interfere with zanamivir binding rarely permit enzyme function. In vitro experiments (86) timed the emergence of mutations conferring resistance to several of the then-new neuraminidase inhibitors. The rapidity with which the virus developed resistance to each compound was directly related to how different the inhibitor molecule was from the structure of the natural substrate. This led to the idea that the closer the drug structure is to that of the natural substrate, the less likely it is that the neuraminidase can mutate and maintain function (87); oseltamivir has a variety of modifications from the natural substrate, and hence resistance was considered more likely to develop to this drug than to zanamivir.

It seems plausible now that both the optimistic predictions for zanamivir and the concerns raised about oseltamivir’s design are being borne out in clinical practice. For zanamivir, no resistant virus has been isolated after treatment of immunocompetent people, but for oseltamivir, the frequency of post-treatment neuraminidase resistance is higher. While only about 0.4% of treated adults harbored influenza viruses with oseltamivir-resistant neuraminidases, this number rose to at least 4% for treated children. Recently, a small study of children in Japan (88) found that, out of 50 children treated with oseltamivir, 9 (18%) harbored viruses with drug-resistance mutations in the neuraminidase gene; the mutations were located where predicted from the in vitro studies discussed above. These mutations occurred far more frequently than has been previously observed (89), but it has not yet been established whether this is a general phenomenon, nor whether the oseltamivir-resistant viruses are transmissible or pathogenic (90).

This tale, both in the elements of resounding success and in the elements of increasing concern about the development of resistance (88), points to the great utility of structural and in vitro studies brought to bear on development of antiviral therapies. In order to continue to benefit from these potent antiviral compounds, it is critical to understand more about which features of neuraminidase inhibitors will discourage the emergence of resistance (90). By understanding the structural basis of resistance, it ought to be possible to design effective neuraminidase inhibitors that are less likely to select for resistant neuraminidase molecules. The same principles should hold true for designing molecules to interact with the sites on the parainfluenza HN protein that participate in receptor binding and F protein triggering.
Using what we know to block parainfluenza virus pathogenesis, and learning more

The multiple roles of the parainfluenza envelope glycoproteins lend themselves to potential strategies for interfering with viral entry, pathogenesis, and disease (Figure 5). Since parainfluenza pathogenesis is likely due in large part to the exuberant inflammatory response to infection, the finding that specific alterations in the HN protein correlate with enhanced pathology, possibly due to the HN protein’s role in induction of inflammatory responses, suggests approaches to modulate this inflammatory response and ameliorate disease (Figure 5A). Given the key role of the inflammatory response as well as the facts that viral replication in the respiratory tract peaks soon after disease onset and that viral titers do not correlate directly with disease severity, any antiviral strategy...
Preliminary studies suggest that it may even be possible to prematurely trigger the F protein, incapacitating it before it can reach its corresponding to the HR repeats of the F protein can be designed specifically to fit into the binding pocket on the globular head of the HN protein (Figure 5B). In addition, this blockade will not only block receptor-mediated fusion (RMF) but also the ability of the HN protein to trigger the F protein, and specific mutations in the stalk region can influence the ability of the HN protein to trigger the F protein, and specific features of the globular head modulate this triggering, it remains completely unknown how the “activating signal” is transmitted from the HN protein to the F protein. If receptor binding induces a conformational change in the HN protein, how is this change communicated to lead to triggering of the F protein? Finally, the HN protein possesses neuraminidase activity and thus the ability to cleave the sialic acid moieties of the cellular receptors, promoting the release of new virions from the host cell surface. Specific inhibition of this activity may prevent virion entry into additional uninfected cells (Figure 5D). These potential therapeutic targets are now being actively pursued and promise to open new avenues for interfering with infection by HPIVs and other viruses.

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