Exploring inhibitor structural features required to engage the 216-loop of human parainfluenza virus type-3 hemagglutinin-neuraminidase†‡

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Human parainfluenza virus type-3 is a leading cause of acute respiratory infection in infants and children. There is currently neither vaccine nor clinically effective treatment for parainfluenza virus infection. Hemagglutinin-neuraminidase glycoprotein is a key protein in viral infection, and its inhibition has been a target for inhibitor development. In this study, we explore the structural features required for Neu2en derivatives to efficiently lock-open the 216-loop of the human parainfluenza virus type-3 hemagglutinin-neuraminidase protein.

Introduction

Human parainfluenza viruses (hPIVs) are a group of respiratory viruses that belong to the Paramyxoviridae family and cause a range of human respiratory diseases in infants, children, immunocompromised patients and elderly people.† Human parainfluenza virus type-3 (hPIV-3) is an enveloped, single-stranded, negative-sense RNA virus that affects children worldwide causing croup, bronchiolitis, and pneumonia.‡ It comes second only to respiratory syncytial virus (RSV) as the leading cause of acute respiratory infections that require hospitalization of infants and children.§ It is also an important cause of morbidity and occasional mortality in immunocompromised patients.‡ Despite continuing efforts, there are currently no effective vaccines or specific therapies to control or treat parainfluenza virus infection.

The envelope of hPIV-3 contains two viral glycoproteins, the hemagglutinin-neuraminidase (HN) and the fusion protein (F).§ The HN protein recognizes sialic acid-containing receptors on respiratory epithelial cell surfaces and uses these sialic acid residues to anchor the virus to the host cell. A second function for the HN protein is promoting F-mediated fusion to allow penetration of the virus genome into the host cell. In addition to its receptor attachment and fusion promotion functions, HN possesses neuraminidase activity that promotes the release of newly formed virions from the cell surface and prevents them from self-agglutination.§,‡ These three functions of the HN protein have made it a major target for anti-hPIV inhibitor design.

Over the past decades, numerous inhibitors that target sialidase proteins of other related viruses, such as influenza A virus (IAV), have been developed. Representative examples are the naturally occurring inhibitor 2-deoxy-2,3-didehydro-N-acetylated neuraminic acid (Neu5Ac2en, 1, Fig. 1)⁷ and the potent anti-IAV 4-deoxy-4-guanidino-Neu5Ac2en inhibitor (zanamivir, 2, Fig. 1).⁸,⁹ However, as a result of the structural differences between hPIV-HN and other sialidases, inhibitory potencies for these unsaturated sialic acid-based inhibitors were very weak against hPIV-HN.¹⁰,¹¹ The first potent Neu2en-based inhibitors to be rationally designed to specifically target hPIV-HN were BCX 2798 (3) and BCX 2855 (4).¹²-¹⁴ In spite of their in vitro potency against hPIV, these two inhibitors failed to continue in development as a result of their poor in vivo efficiency.¹²-¹⁴ We have recently reported¹¹ the development of potent Neu2en-based inhibitors (5 and 6) that carry bulky C-4 substituents. These bulky C-4 substituents target a

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Fig. 1 Structures of the anti-influenza sialidase inhibitors (1 and 2), the hPIV inhibitors (3 and 4), and the potent hPIV-3 designer inhibitors (5 and 6).
unique feature in the hPIV-3 HN protein, the flexible 216-loop, locking open the loop and efficiently occupying the created 216-cavity.\textsuperscript{11} This efficient binding to the HN protein results in potent inhibition of both HN functions, as well as significant blockade of virus propagation.

The ability of these new designer inhibitors (5 and 6) to force the 216-loop open and reveal the 216-cavity is attributed to their bulky C-4 functionality (the 4-phenyl-1,2,3-triazole group). This outcome has proved the notion that the binding pocket of the hPIV-3 HN protein can efficiently accommodate Neu2en derivatives carrying bulky, hydrophobic C-4 substituents.\textsuperscript{15} Such bulky inhibitors are not only able to fit into the binding pocket but also benefit from the extra hydrophobic binding interactions exerted by their C-4 bulky substituents, resulting finally in more potent inhibition of HN functions.

The STD NMR experiments carried out for these triazole inhibitors in complex with either the intact virus or the recombinant HN protein have shown that the most significant interactions with the protein were exerted by the phenyl moiety carried by the 1,2,3-triazole ring.\textsuperscript{11} Accordingly, in this study, we verify the importance of the triazole ring as a spacer carrying the hydrophobic phenyl group at the C-4 position of 4-deoxy-Neu2en, by replacing it with other acyclic spacers (urea,\textsuperscript{16} amide and sulfonamide functionalities), in order to explore the structural features of the C-4 substituent and define the essential inhibitor characteristics required to force the 216-loop open to reveal and efficiently occupy the 216-cavity.

**Results and discussion**

**Computational chemistry**\textsuperscript{17}

In the lead inhibitors (5 and 6), the distance between the triazole N-1 and C-1 of the phenyl ring in the energy minimized structures was found to be 3.58 Å. Accordingly, the triazole ring was replaced by a number of acyclic spacers that hold the phenyl ring at comparable distances from the 4-deoxy-Neu5Ac2en skeleton. The selected spacers, which can be readily synthesized using common 4-deoxy-Neu5Ac2en intermediates, are urea-, amide- and sulfonamide-based. These three linkers were found to generate distances of 3.65 Å, 2.44 Å and 2.87 Å, respectively, between the 4-amino-4-deoxy-Neu5Ac2en C-4 nitrogen and C-1 of the phenyl ring (Fig. 2).

As predicted from these preliminary calculations and by aligning the structures of these 4-deoxy-Neu5Ac2en derivatives that contain these three acyclic linkers, the urea-based linker creates the most comparable distance between the 4-deoxy-Neu5Ac2en skeleton and the phenyl ring in the triazole inhibitor 5 (Fig. 3).

**Chemistry**

Based on the initial computational chemistry calculations, a series of 4-deoxy-4-phenylureido-Neu5Ac2en-based inhibitors that contain variable substituents at the phenyl group (9a–g, Scheme 1) was synthesized. The synthesis of these urea derivatives was achieved by using the known intermediate 7.\textsuperscript{10} As shown in Scheme 1, the fully protected 4-amino-4-deoxy-Neu5Ac2en intermediate (7) was reacted with a number of substituted phenyl and benzyl isocyanate reagents in dichloromethane, in the presence of a catalytic amount of DMAP, to yield the protected urea derivatives 8a–g in excellent yields. The deprotection of compounds 8a–g by treatment with NaOH in MeOH:H\textsubscript{2}O (1:1), provided the deprotected target 4-deoxy-4-arylureido-Neu5Ac2en derivatives 9a–g.

The same intermediate 7 was utilized (Scheme 2) for the synthesis of amides 11a–c by reaction with three different benzoyl chlorides in the presence of triethylamine to yield the fully protected amides 10a–c, which upon stirring in a 1:1 mixture of methanol:water at pH 13–14 (adjusted by NaOH) afforded the final deprotected amides 11a–c.

In Scheme 3, amine 7 was converted to the protected sulfonamides 12a–d by reaction with the appropriate sulfonyl chloride in the presence of triethylamine. The resultant sulfonamides 12a–d were then deprotected by treatment with NaOH in 50% MeOH to afford the target sulfonamides 13a–d.

In order to evaluate the effect of a C-5 modification on the inhibitory potency of these derivatives, the reported 4-amino-4-deoxy-3-isobutylamido-Neu2en intermediate (14) (ref. 10) was utilized to synthesize amide 15 as shown in Scheme 4.
Thus, stirring amine 14 in the presence of triethylamine with 4-chlorobenzoyl chloride afforded amide 15 in 85% yield. Deprotection of the resultant amide by exposure to NaOH in aq. MeOH yielded the final deprotected inhibitor 16 in 94% yield.

Biological screening

Evaluation of the synthesized urea-, amide- and sulfonamide-based 4-deoxy-Neu2en derivatives for their capacity to block the neuraminidase function of hPIV-3 HN, as well as a comparison of their potencies to those of the lead triazole inhibitors 5 and 6, was carried out by using a standard neuraminidase inhibition (NI) assay (Fig. 4).11

Surprisingly, and in spite of the close spacer length measurements between the triazole ring and the urea functionalities, all of the synthesized C-4 ureido-Neu5Ac2en derivatives failed to show NI IC50 values below the maximal-threshold concentration used in this assay (150 μM, data not shown). On the other hand, the neuraminidase inhibition activities exerted by amide and sulfonamide derivatives were relatively more potent and comparable to each other. Within each of these two series (amides and sulfonamides), the most potent neuraminidase inhibition activity was exerted by the 4-chlorophenyl derivatives (11b and 13b), with the amide 11b showing an IC50 value of 45.1 μM and the sulfonamide 13b showing an IC50 value of 48.7 μM. By comparing the three amides, the next potent neuraminidase inhibitor within this group was found to be the phenylamido derivative 11a with an IC50 value of 103.5 μM, while derivative 11c with a 4-methoxyphenylamido group failed to show an IC50 value below the detection threshold. Within the sulfonamide series, the 4-toluenesulfonamido derivative 13d was a poorer inhibitor compared to the 4-chlorophenylsulfonamide 13b with an IC50 value of 64.8 μM. The 4-methoxyphenylsulfonamide 13c was found to be slightly less potent with an IC50 of 71.0 μM, while the weakest neuraminidase inhibitor within the
sulfonamides was the phenylsulfonamido derivative 13a with an IC$_{50}$ of 99.6 μM.

The unexpected weak inhibition observed for all of the urea derivatives can be rationalized on the basis of the significant flexibility of the phenyl moiety-bearing urea linker in these derivatives, in contrast to the rigid triazole ring in the potent inhibitors 5 and 6. It is reasonable to conclude from these results that the linker that carries the bulky phenyl substituent at the C-4 position of Neu2en needs to be rigid enough to keep the phenyl ring maximally oriented towards the 216-loop and occupy the open loop 216-cavity. In both the amide and the sulfonamide derivatives, the linker holds the phenyl group at a shorter distance relative to that in the C-4 urea derivatives. Moreover, these linkers are less flexible and direct the C-4 phenyl moiety towards the open loop 216-cavity to provide the observed improved potency within these two series compared with the C-4 urea derivatives. Conversely, in the inhibitor series 11 and 13 the combination of C-4 phenyl moieties positioned at a shorter distance from the 216 loop with the use of more flexible linkers, compared to the rigid triazole linker in inhibitors 5 and 6, resulted in a decreased potency. This combination presumably limits the inhibitors’ capability to lock open the 216-loop and efficiently occupy the 216-cavity.

In order to evaluate the effect of C-5 substitution in these new inhibitors on their NI potencies, the C-5 acetamido group of the derivative that showed the most potent inhibition (the 4-chlorophenylamide 11b) was replaced by the isobutyramido group that has been reported in previous studies to further improve the inhibition of Neu2en derivatives against hPIV-3 HN. Accordingly, inhibitor 16 that contains a C-4 4-chlorophenylamido substituent and a C-5 isobutyramido group was synthesized and tested for its capacity to inhibit hPIV-3 HN neuraminidase activity. Indeed, the new C-5 isobutyramido-based inhibitor proved to be slightly more potent with a NI IC$_{50}$ value of 38.9 μM (vs. 45.1 μM for the parent C-5 acetamido derivative 11b), suggesting a similar binding mode to that of the triazole derivatives.

Conclusions

In conclusion, a group of C-4 phenylureido, phenylamido and phenylsulfonamido 4-deoxy-Neu2en derivatives, in which the triazole linker carrying the phenyl moiety in inhibitors 5 and 6 was replaced by more flexible linkers, have been synthesized and evaluated for their capacity to inhibit hPIV-3 HN. The results have shown that the triazole ring in inhibitors 5 and 6 plays an important role in potency. Indeed, the rigidity of the triazole ring is critical for inhibitor potency, as it suitably orients the phenyl ring towards the 216-loop. This optimal linker can induce 216-loop movement to reveal the 216-cavity, allowing the inhibitor to efficiently occupy the cavity and tightly bind HN. Importantly, the more flexible the linker carrying the phenyl ring is, the less capable the inhibitor is to induce the 216-loop movement, resulting in weaker inhibition of hPIV-3 HN functions. This was illustrated by the NI IC$_{50}$ values of the urea, amide and sulfonamide derivatives evaluated in this study, where the weakest inhibition was observed for the most flexible derivatives (the urea series), while intermediate inhibition was found for the derivatives that have linkers with moderate rigidity (the amides and sulfonamides). These results set the basis for the design of novel Neu2en derivatives carrying bulky C-4 substituents targeting hPIV-3 HN with the aim of reorienting the 216-loop and occupying the 216-cavity.

Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. I. M. E.-D. and P. G. contributed equally and are joint senior authors.

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Notes and references


17 Computational Chemistry calculations were carried out using ‘Molecular Operating Environment (MOE) version 2008.10’, Chemical Computing Group Inc., 1010 Sherbrooke Street West, Suite 910, Montreal, H3A 2R7, Canada.