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A multi-pronged evaluation of aldehyde-based tripeptidyl main protease inhibitors as SARS-CoV-2 antivirals

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** Abbreviations: COVID-19, coronavirus disease 2019; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2; MPro, main Protease; MPI, main Protease Inhibitor; SAR, structure-activity relationship; eGFP, enhanced green fluorescent protein; CBZ, benzylxycarbonyl.

A multi-pronged evaluation of aldehyde-based tripeptidyl main protease inhibitors as SARS-CoV-2 antivirals

As an essential enzyme of SARS-CoV-2, the COVID-19 pathogen, main protease (M\textsubscript{Pro}) is a viable target to develop antivirals for the treatment of COVID-19. By varying chemical compositions at both P2 and P3 positions and the N-terminal protection group, we synthesized 18 tripeptidyl M\textsubscript{Pro} inhibitors that contained also an aldehyde warhead and \(\beta\)-(S-2-oxopyrrolidin-3-yl)-alaninal at the P1 position. Systematic characterizations of these inhibitors were conducted, including their \textit{in vitro} enzymatic inhibition potency, X-ray crystal structures of their complexes with M\textsubscript{Pro}, their inhibition of M\textsubscript{Pro} transiently expressed in 293T cells, and cellular toxicity and SARS-CoV-2 antiviral potency of selected inhibitors. These inhibitors have a large variation of determined \textit{in vitro} enzymatic inhibition IC\textsubscript{50} values that range from 4.8 to 650 nM. The determined \textit{in vitro} enzymatic inhibition IC\textsubscript{50} values reveal that relatively small side chains at both P2 and P3 positions are favorable for achieving high \textit{in vitro} M\textsubscript{Pro} inhibition potency, the P3 position is tolerable toward unnatural amino acids with two alkyl substituents on the \(\alpha\)-carbon, and the inhibition potency is sensitive toward the N-terminal protection group. X-ray crystal structures of M\textsubscript{Pro} bound with 16 inhibitors were determined. In all structures, the M\textsubscript{Pro} active site cysteine interacts covalently with the aldehyde warhead of the bound inhibitor to form a hemithioacetal that takes an S configuration. For all inhibitors, electron density around the N-terminal protection group is weak indicating possible flexible binding of this group to M\textsubscript{Pro}. In M\textsubscript{Pro}, large structural variations were observed on residues N142 and Q189. Unlike their high \textit{in vitro} enzymatic inhibition potency, most inhibitors showed low potency to inhibit M\textsubscript{Pro} that was transiently expressed in 293T cells. Inhibitors that showed high potency to inhibit M\textsubscript{Pro} transiently expressed in 293T cells all contain \(\alpha\)-tert-butyl-threonine at the P3 position. These inhibitors also exhibited relatively low cytotoxicity and high antiviral potency. Overall, our current and previous studies indicate that \(\alpha\)-tert-butyl-threonine at the P3 site is a key component to achieve high cellular and antiviral potency for tripeptidyl aldehyde inhibitors of M\textsubscript{Pro}. 

\textit{Abbreviations}: COVID-19, coronavirus disease 2019; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2; MPro, main Protease; MPI, main Protease Inhibitor; SAR, structure-activity relationship; eGFP, enhanced green fluorescent protein; CBZ, benzylxycarbonyl.
1. Introduction

COVID-19 is the current prevailing pandemic that has devastated much of the world. As the COVID-19 pathogen, SARS-CoV-2 uses its membrane Spike protein to recognize the human receptor ACE2 for infection [1,2]. Current COVID-19 vaccines all target this process for the neutralization of the virus. However, the continuous emergence of new viral strains that evade vaccines demands other antivirals to be developed as well. The genome of SARS-CoV-2 encodes a large open reading frame ORF1ab that is translated to two large polypeptides pp1a and pp1b in the human cell host [3,4]. The processing of pp1a and pp1b to 16 nonstructural proteins (nsps) that are functionally critical to the viral replication relies on proteolytic activities of two internal nsp fragments, nsp3 and nsp5 [5]. Nsp5 is also called 3C-like protease and more recently main protease (MPro). Since MPro hydrolyzes 13 out of the total of 16 nsps, it has been considered as a viable target for the development of antivirals. In the past year, a number of papers have been published on the development of peptidyl aldehydes that contain β-(S-2-oxopyrrolidin-3-yl)-alaninal (opal) at the P1 position as potent MPro inhibitors [6-14]. Other inhibitors were developed as well including nirmatrelvir that has been approved by the U.S. Food and Drug Administration for its combined use with ritonavir as a therapy, branded as pxlovid, for COVID-19 and S-217622 that is undergoing clinical trials for COVID-19 in Japan [15-30]. However, systematic structure-activity relationship (SAR) studies of opal-based MPro inhibitors are needed. In the current study, we explore variations at the P2 and P3 positions and the N-terminal protection group in opal-based tripeptidyl MPro inhibitors. Our study indicates that O-tert-butylenine at the P3 position in an opal-based MPro inhibitor is critical for achieving high potency inhibiting MPro expressed in human host cells and antiviral potency against SARS-CoV-2.

Fig. 1. Structures of tripeptidyl MPro inhibitors MPI11–28. Inhibitors that have just a single component at R2/R3 shown contain a hydrogen at the R3 position. MPI13-8 were previously developed and are shown for comparison.
2. Results

2.1. The design and synthesis of MPI11-28

In previous studies, we developed MPI1-10 and characterized both their MPro inhibition potency and their antiviral potency [13,14]. As a tripeptidyl aldehyde, MPI8 shows the highest antiviral potency with an IC50 value of 30 nM to neutralize SARS-CoV-2 (USA-WA1/2020) in Vero E6 cells [31]. MPI3, another tripeptidyl aldehyde, doesn’t have high antiviral potency. However, it has the highest in vitro MPro inhibition potency with an IC50 value of 8.5 nM [13]. To explore how substrituents at different positions in a tripeptidyl opal-based MPro inhibitor (Fig. 1) influence its potency, we decided to carry out a systematic structure-activity relationship (SAR) study. We maintained opal at the P1 position due to its established preferential binding to the MPro P1 binding pocket and its ability to form a covalent adduct with C145, the MPro catalytic cysteine. MPro tolerates leucine and phenylalanine at the P2 position in a substrate. Therefore, we chose to vary this site in our inhibitor design among β-alkyl alamines with sizes between or around leucine and phenylalanine. Chosen alkyl substituents include isopropyl, phenyl, cyclohexyl, t-butyl, isopropenyl, cyclopropyl, 2-furyl, and 2-thienyl. MPro doesn’t have a binding pocket for the P3 position in a substrate. However, native MPro substrates have valine or a similar size residue at this position. Based on known opal inhibitors and substrates of MPro, we varied this site among amino acids including valine, O-tert-butyl-threonine, L-cyclopentylglycine, L-tert-butyl-glycine, L-α-methyl-valine, dimethylglycine, and 1-aminoxycholepropane-1-carboxylate. The N-terminal protection group was chosen among benzylloxycarbonyl (CBZ), m-chloro CBZ, and indole-2-carboxylate due to their demonstrated contributions to antiviral potency [6,10]. A total of 18 new MPro inhibitors, designated as MPI11-28 shown in Fig. 1, were designed. We synthesized all inhibitors according to the synthetic route shown in Scheme 1. In this synthesis, a P2 amino acid ester was conjugated with a N-protected P3 amino acid and the afforded product was then hydrolyzed to free its C-terminal carboxylate for a reaction with β-(S-2-oxopyrrolidin-3-yl)-alanine ester. An obtained tripeptidyl ester was reduced to afford a β-(S-2-oxopyrrolidin-3-yl)-alanine ester. An obtained tripeptidyl ester was then hydrolyzed to free its β-C-terminal carboxylate for a reaction with Dess-Martin oxidation in a mild condition to make a final product.

2.2. Kinetic characterizations of MPI11-28 on their enzymatic inhibition of MPro

We followed a previously established protocol that uses Sub3 (Dabcsyl-KTSALQSGFKME-Edans), a fluorogenic peptide substrate of MPro to determine IC50 values for MPI11-28 [28]. In this assay, we preincubated MPro with an inhibitor for 30 min before Sub3 was added and the fluorescent product formation (Ex: 336 nm/Emission: 455 nm) was recorded in a fluorescence plate reader. 30 min incubation time with MPro is a standard procedure that has been used by multiple labs in the determination of IC50 values for MPro inhibitors [11,27,28,32]. Since MPI11-28 are reversible covalent inhibitors, their incubation times with MPro are not expected to significantly influence their determined IC50 values. Tests involving MPI11 in three different incubation times, 15, 30, and 60 min, led to very similar determined IC50 values for all inhibitors. As shown in Table 1, MPI11-28 display a large variation of IC50 values that ranges from 4.8 to 650 nM. MPI11-14 all contain valine at the P3 position. They are among the most in vitro potent inhibitors for MPro. MPI13-14 have determined IC50 values around 5 nM. Since 10 nM MPro is the lowest enzyme concentration we can use to do the inhibition analysis, 5 nM is technically the lowest IC50 value we can detect. Therefore, MPro inhibition potency for MPI13-14 is likely higher than the numbers shown in Table 1. In comparison to MPI13-14, MPI11-12 have slightly higher IC50 values. In previous work, we developed another three MPro inhibitors MPI3-5 that also contain a valine at the P3 site. All three have low IC50 values. All 7 inhibitors that have valine at the P3 position and a N-terminal CBZ protection group are among the most potent MPro inhibitors in vitro. A P3 Valine apparently favors the enzyme inhibition kinetics. A comparison of all seven inhibitors also reveals that a residue at the P2 position with a size close to leucine leads to better MPro inhibition. β-Cyclopentyl alanine and β-(furan-2-yl) alanine that have a more rigid side chain than leucine favor the enzyme inhibition kinetics more. MPI6-8 that have a P3 O-tert-butyl-threonine were developed previously and showed high antiviral potency. We synthesized two more MPro inhibitors with a P3 O-tert-butyl-threonine, MPI16-17. Both have an IC50 value similar to MPI6-8. Varying the P2 position does not seem to significantly change in vitro MPro inhibition potency among this group of inhibitors. To explore further on whether variations at the P3 position lead to different inhibition potency, we developed MPI18-24. Variants include two dialkyl glycines and 1-aminoctylpropene-1-carboxylate that are not standard β-amino acids. Except MPI23 that contains a P3 1-aminocyclopropane-1-carboxylate, all other inhibitors have an IC50 value below 100 nM. The more rigid P3 position in MPI23 likely contributes to its less favorable MPro inhibition kinetics. MPI24 is structurally similar to MPI13. Its enzyme inhibition potency is similar to MPI13 as well. Three inhibitors MPI25-27 that have a N-terminal m-chloro CBZ group were developed and characterized as well. In comparison to their regular CBZ-containing counterpart inhibitors MPI8, MPI6, and MPI5, they display more than 5-fold higher IC50 values. Apparently adding an m-chloro substituent to the terminal CBZ group doesn’t improve in vitro...
enzyme inhibition potency. MPI28 has an N-terminal indole-2-carboxylate. In comparison to its CBZ counterpart inhibitor MPI3, MPI28 has a 6-fold higher IC50 value. In comparison to indole-2-carboxylate, it is evident that CBZ serves as a better N-terminal group for improved in vitro enzyme inhibition potency.

2.3. X-ray crystallography analysis of MPro bound with 16 different inhibitors

To characterize our developed inhibitors on their interactions with MPro, we crystallized MPro in its apo form, soaked apo-MPro crystals with different inhibitors and then determined structures of MPro bound with different inhibitors using X-ray crystallography. Among 17 inhibitors...
that we used to soak crystals, 16 were observable in the active site of M\textsuperscript{Pro}. All structures were determined to high resolutions (Table S1). For all inhibitors, their calculated 2F\textsubscript{o}-F\textsubscript{c} electron density maps in the M\textsuperscript{Pro} active site clearly show a covalent interaction with M\textsuperscript{Pro} C145 to generate a hemithioacetal (Fig. 3A). In all formed hemithioacetals, the hemithioacetal alcohol takes an \textit{S} configuration. In most calculated structures, electron density around inhibitors have defined shapes for modeling the P1, P2 and P3 residues. For MPI12, the isopropenyl group has very weak electron density around its C1 and C3 atoms, indicating that the isopropenyl group is probably freely rotating around its C2 atom in the active site of M\textsuperscript{Pro}. For MPI26 and MPI28 that have a P2 leucine, electron density around the side chain isopropyl group is also weak, indicating its flexible conformations at the M\textsuperscript{Pro} active site. Although the N-terminal group of inhibitors including MPI11, MPI12, MPI18, and MPI19 were successfully modeled, most calculated structures have relatively weak electron density around the inhibitor N-terminal group, indicating weak binding of the N-terminal group at the active site.

In M\textsuperscript{Pro}-MPI11 as shown in Fig. 3B, MPI11 forms extensive hydrogen bonds with M\textsuperscript{Pro}. The hemithioacetal hydroxyl group is within the hydrogen bond distance to three backbone \textalpha-amines in residues 143–145 that form an oxyanion hole to facilitate the enzyme-catalyzed hydrolysis of a substrate. Due to the preferential binding of a negative charged hydroxyl group at this position, it is likely that the hemithioacetal hydroxy group is deprotonated to strongly interact with the three backbone \textalpha-amines, explaining the exclusive \textit{S} configuration of the formed hemithioacetal. The opal side-chain lactam forms three hydrogen bonds with residues E166, N142, and F140 and its backbone \textalpha-amine forms a hydrogen bond with H164. The side chain of the P2 \beta-tert-butyl-alanine (4-methyl-leucine) is involved in van der Waals interactions with M\textsuperscript{Pro} residues including M49, H41, M165, D187, and

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**Fig. 3.** Crystal structures of M\textsuperscript{Pro} bound with 16 MPIs. (A) Contoured 2F\textsubscript{o}-F\textsubscript{c} maps at the 1\textsigma level around 16 MPIs and C145 in the active site of M\textsuperscript{Pro}. (B–Q) Interactions between 16 MPIs and the M\textsuperscript{Pro} active site residues.
Q189. It is interesting to note that the M49-containing loop has weak electron density around it, indicating flexible conformations of this loop when MPI11 binds to the $\text{M}^{\text{Pro}}$ active site. The MP11 P2 $\alpha$-amine interacts with the side chain of Q189 indirectly through two water-bridged hydrogen bonds. The $\text{M}^{\text{Pro}}$-MPI11 structure shows that the tert-butyl group is within a close van der Waals distance to the Q189-containing loop of $\text{M}^{\text{Pro}}$. To accommodate a chemical moiety bigger than $\beta$-tert-butyl-alanine, M$^{\text{Pro}}$ will need to rearrange the Q189-containing loop to enlarge the P2 binding pocket. The P3 valine of MPI11 forms two hydrogen bonds with the backbone $\alpha$-amine and $\alpha$-carbonyl groups of E166. Its side chain forms van der Waals interactions with the side chain of E166. The N-terminal CBZ of MPI11 interacts hydrophobically with the backbone of the Q189-containing loop and the side chain of P168. However, its weak electron density indicates loose binding. In M$^{\text{Pro}}$-MPI11, the side chain of N142 flips away from the opal side chain of MPI11. The $\text{M}^{\text{Pro}}$-MPI12 structure around the active site is shown in Fig. 3D. Interactions between MPI13 and M$^{\text{Pro}}$ are mostly similar to that between MPI12 and M$^{\text{Pro}}$. N142 adopts a conformation like the one in $\text{M}^{\text{Pro}}$-MPI11. Unlike MPI12, the P2 cyclopropyl side chain of MPI13 has defined, strong electron density around it and interacts with the van der Waals distance with side chains from H41, D187 and Q189. This observation explains high in vitro enzymatic inhibition potency for inhibitors including MPI13, MPI19 and MPI24. MPI14 shown in Fig. 3E interacts with M$^{\text{Pro}}$ mostly similar to MPI11. MPI14 has a P2 $\beta$-(2-furanyl)-alanine residue that has defined, strong electron density around the furan side chain. The furan group is in parallel with the H41 imidazole side chain so that the two aromatic groups are likely engaged in a $\pi$-$\pi$ stacking interaction. This $\pi$-$\pi$ stacking interaction likely contributes to the high in vitro enzymatic inhibitor potency of MPI14. In M$^{\text{Pro}}$-MPI14, Q189 has weak electron density around its side chain. It is not involved in hydrogen bonding interactions with MPI14 directly or indirectly. In both M$^{\text{Pro}}$-MPI13 and M$^{\text{Pro}}$-MPI14, the M49-containing loop has weak electron density. MPI16 differs from MPI11 in its P3 position that is O-tert-butyl-threonine. As shown in Fig. 3F, MPI16 maintains most hydrogen bonding interactions that show in the $\text{M}^{\text{Pro}}$-MPI11 complex structure. The $\text{M}^{\text{Pro}}$ Q189 side chain amide is 3.5 Å to the $\alpha$-amine of MPI16. This is around the borderline for a hydrogen bond interaction. In $\text{M}^{\text{Pro}}$-MPI16, the N142 side chain adopts a conformation that caps the MPI16 P1 opal side chain. Compared to MPI11, MPI16 has an additional P3 O-tert-butyl-group. Its location can be clearly refined that shows van der Waals interactions with P168 and E166. MPI18 has a chemical structure very similar to MPI11 except at the P3 position that has an additional methyl group on the $\beta$-carbon. Its interactions with M$^{\text{Pro}}$ as shown in Fig. 3G are also similar to that involving MPI11. Visible differences are at M$^{\text{Pro}}$ residues N142 and Q189. The N142 side chain adopts a conformation to interact on the M188 opal side chain of MPI18. The Q189 side chain amide forms a hydrogen bond with the N-terminal carbamoyl oxygen atom. MPI19 is structurally similar to MPI13 except an additional methyl group at the P2 $\beta$-carbon position. As shown in Fig. 3H, MPI19 interacts with M$^{\text{Pro}}$ similar to MPI13 except that the N142 side chain in $\text{M}^{\text{Pro}}$-MPI19 adopts a conformation to cap the M$^{\text{Pro}}$ P1 binding pocket for the MPI19 P1 opal residue. Except a cyclopropyl group at the P3 position, MPI20 is structurally similar to MPI11. As shown in Fig. 3I, MPI20 interacts with M$^{\text{Pro}}$ similar to MPI11. A water molecule bridges the interaction between the Q189 side chain amide and the MPI20 P2 $\alpha$-amine. The N142 side chain caps the M$^{\text{Pro}}$ P1 binding pocket. In M$^{\text{Pro}}$-MPI20, electron density around residues 45–50 is too weak to model these residues, indicating that this loop region adopts a highly flexible conformation. The same region shows weak electron density in M$^{\text{Pro}}$-MPI16, M$^{\text{Pro}}$-MPI18 and M$^{\text{Pro}}$-MPI19. MPI21 has $\alpha$-methyl-valine at its P3 position. Compared to MPI11, the additional P3 $\alpha$-methyl group pushed the N-terminal carbamate of MPI21 to adopt an orientation that rotates about 90° from the one shown in the M$^{\text{Pro}}$-MPI11 (Fig. 3J). Although the Q189 side chain amide is within the hydrogen bond distance to the MPI21 N-terminal carbamate carbonyl oxygen atom, the geometry is not right. Compared to other inhibitors that have a regular $\alpha$-amino acid at their P3 position, this different orientation at the N-terminal carbamate in MPI21 apparently leaves space for the Q189 side chain to move deeper toward the M$^{\text{Pro}}$ active site. In M$^{\text{Pro}}$-MPI21, the N142 side chain adopts a conformation similar to the one shown in M$^{\text{Pro}}$-MPI11. MPI21 is structurally similar to MPI21 but contains dimethylglycine at its P3 position. The $\text{M}^{\text{Pro}}$-MPI22 structure at the M$^{\text{Pro}}$ active site is very similar to M$^{\text{Pro}}$-MPI12. The pro-S-methyl group at the MPI22 P3 residue pushes the N-terminal carbamate to adopt a conformation that positions its carbamate carbonyl oxygen away from forming a hydrogen bond with the Q189 side chain amide (Fig. 3K). As same as in M$^{\text{Pro}}$-MPI21, this new orientation at the N-terminal carbamate leaves space for the Q189 side chain to move deeper toward the M$^{\text{Pro}}$ active site. MPI23 has a cyclopropyl backbone at its P3 position. Similar to MPI21 and MPI22, this cyclopropyl group pushes the N-terminal carbamate to adopt a conformation that positions the carbamate carbonyl oxygen atom away from forming a hydrogen bond with the Q189 side chain amide through the rotating extend of the carbamate is not as much as in MPI21 and MPI22 (Figure 3L). The P2 position of MPI23 is $\beta$-cyclohexyl-alanine. The cyclohexyl group has defined, strong density that indicates a chair conformation. The cyclohexyl group aligns parallelly with respect to the H41 side chain, making two C-H groups from the cyclohexyl group directly interact with the $\pi$ orbital of the H41 side chain. Compared to MPI11, the P2 residue of MPI23 is larger and pushes the Q189-containing loop to move slightly away from the active site. MPI24 has a structure similar to MPI13. It has a cyclopropyl group compared to an isopropyl group in MPI13 at the P3 $\beta$-carbon. The interactions between MPI24 and M$^{\text{Pro}}$ are also similar to that between MPI13 and M$^{\text{Pro}}$ (Figure 3M). In M$^{\text{Pro}}$-MPI24, the N142 side chain caps the M$^{\text{Pro}}$ P1 binding pocket and the Q189 side chain amide forms a hydrogen bond with the MPI24 P2 $\alpha$-amine. In M$^{\text{Pro}}$-MPI21, M$^{\text{Pro}}$-MPI22, M$^{\text{Pro}}$-MPI23, and M$^{\text{Pro}}$-MPI24, the M49-containing loop has weak electron density that leads to structures of certain residues in M$^{\text{Pro}}$-MPI21 and M$^{\text{Pro}}$-MPI22 not able to be refined. MPI25 is a P18 derivative. Its interactions with M$^{\text{Pro}}$ involve most hydrogen bonds that exist in the $\text{M}^{\text{Pro}}$-MPI11 structure (Figure 3N). In M$^{\text{Pro}}$-MPI25, the N142 side chain points away from the position that can cap the M$^{\text{Pro}}$ P1 binding pocket and the Q189 side chain amide interacts with the MPI25 P3 side chain oxygen through two water-bridged hydrogen bonds. Since the electron density around the N-terminal group is weak, the chloride position cannot be unambiguously assigned but it is with high confidence that it interacts with the P168 side chain. In M$^{\text{Pro}}$-MPI25, the M49-containing loop also has relatively weak electron density so that the M49 side chain conformation cannot be unambiguously assigned either. The O-tert-butyl group in the P3 residue has strong electron density that was used to unambiguously refine its structure. The P168 and E166 side chains of MPI26 is structurally different from MPI25 only at the P2 residue. As shown in Figure 3O, the binding model of MPI26 at the M$^{\text{Pro}}$ active site is very similar to MPI23. Visible differences are at M$^{\text{Pro}}$ residues N142 and Q189. In M$^{\text{Pro}}$-MPI26, the N142 side chain adopts a conformation that caps the M$^{\text{Pro}}$ P1 binding pocket and the Q189 side chain amide forms hydrogen bonds separately with the P162 P2 $\alpha$-amine and the N-terminal carbamate carbonyl oxygen. The electron density around residues 45–50 is also too weak to be used to refine their structures. The electron density around the side chains of P168 and E166 of MPI26 is very similar to that of MPI25.
density around the P2 leucine β-isopropyl group allows only one methyl group can be clearly refined, indicating that the P2 leucine β-isopropyl group may adopt different conformations. MPI27 is structurally different from MPI25 only at its P3 position that is valine. As shown in Figure 3P, MPI27 interacts with M\textsuperscript{Pro} different from MPI25 only at its P3 position that is valine. As shown in Figure 3P, MPI27 interacts with M\textsuperscript{Pro} similar to MPI25. In M\textsuperscript{Pro}-MPI27, the P3 valine side chain packs on top of the E166 side chain to form van der Waals interactions. The Q189 side chain amide forms a hydrogen bond with the MPI27 P2 α-amide. Unlike all other inhibitors, MPI28 has a N-terminal indole-2-carbonyl protection group. Its interactions with M\textsuperscript{Pro} shown in Figure 3Q are similar to that in M\textsuperscript{Pro}-MPI11. The Q189 side chain amide forms a hydrogen bond with the N-terminal amide oxygen. As in most other M\textsuperscript{Pro}-inhibitor structures, the electron density at the N-terminal group is too weak to refine its accurate structure. Like in M\textsuperscript{Pro}-MPI26, the electron density at the P2 leucine β-isopropyl group allows the refinement of only one methyl group, indicating flexible binding of this group at the M\textsuperscript{Pro} active site. In M\textsuperscript{Pro}-MPI28, the electron density around M\textsuperscript{Pro} residues 46–50 is too weak to be used to refine their structures. Although some of M\textsuperscript{Pro}-inhibitor complexes have refined structures for residues 45–50, all of them have relatively weaker electron density around these residues than other parts of M\textsuperscript{Pro}, indicating that the M49-containing loop is flexible after an inhibitor binds to the M\textsuperscript{Pro} active site.

2.4. Characterizations of cellular M\textsuperscript{Pro} inhibition potency of MPI11-28

M\textsuperscript{Pro} is acutely toxic to a human cell host. Using this unique characteristic, we developed a cell-based analysis to characterize cellular M\textsuperscript{Pro} inhibition potency for compounds [31]. A similar assay was also used by others to analyze cellular potency of M\textsuperscript{Pro} inhibitors [33,34]. In this assay, a compound with cellular potency inhibits cellular toxicity from the expression of an M\textsuperscript{Pro}-eGFP fusion protein, leading to enhanced overall expression of M\textsuperscript{Pro}-eGFP that can be characterized by flow cytometry. This assay is more advantageous over an antiviral assay in analyzing the cellular M\textsuperscript{Pro} inhibition ability of a compound since it may inhibit host proteases such as TMPRSS2, furin, and cathepsin L that are critical for SARS-CoV-2 infection to provide false positive cellular potency results for M\textsuperscript{Pro} inhibition [35–37]. We used this method to characterize a number of investigational M\textsuperscript{Pro} inhibitors. Our study showed some inhibitors that have demonstrated antiviral potency do not inhibit M\textsuperscript{Pro} directly in a human cell host, indicating different mechanisms for these compounds to achieve antiviral potency [31]. Other advantages of using this assay include quickly priming potent inhibitors for their antiviral assays by bypassing cellular permeability and stability tests and avoiding time-consuming and more complicated antiviral tests that involve the use of a biosafety level 3 facility for compounds with low cellular potency. We adopted this assay to characterize MPI11-28. All inhibitors were tested up to 10 μM in their inhibition of M\textsuperscript{Pro}-eGFP that was transiently expressed in human 293T cells. Overall cellular eGFP fluorescence was plotted against the inhibition concentration to obtain their EC\textsubscript{50} values. Results are presented in Fig. 4 and Table 1. As shown in Fig. 4A, MPI11-14 exhibited minimal cellular potency to inhibit M\textsuperscript{Pro} in 293T cells, which is in significant contrast to their very high in vitro enzyme inhibition potency. Since their inhibition curves do not reach a plateau, their EC\textsubscript{50} values are estimated as > 2 μM. MPI16 and MPI17 are the two most potent inhibitors among MPI11-28 on cellular potency with determined EC\textsubscript{50} values as 56 nM and 97 nM, respectively (Fig. 4B). In comparison to MPI16 and MPI17, MPI18 and MPI19 have weaker cellular potency with determined EC\textsubscript{50} values as 860 nM and 1500 nM, respectively. MPI20-23 all have weak cellular potency (Fig. 4C). MPI22 and MPI23 showed very low inhibition of M\textsuperscript{Pro}-eGFP in 293T cells. At 10 μM for these two compounds, M\textsuperscript{Pro}-eGFP expression was lower than half of the plateau level of M\textsuperscript{Pro}-eGFP that was observed for MPI16-19. For these four inhibitors, their EC\textsubscript{50} values are estimated as > 2 μM for MPI20 and MPI21 and > 10 μM for MPI22 and MPI23. Among MPI24-28 (Fig. 4D), MPI25-27 exhibited high cellular potency with EC\textsubscript{50} values as 140 nM, 230 nM, and 630 nM respectively. On contrary to their high in vitro enzyme inhibition potency, MPI24 and MPI28 have very weak cellular potency with estimated EC\textsubscript{50} values as > 2 μM and >10 μM, respectively. Two reported M\textsuperscript{Pro} inhibitors nirmatrelvir [29] and GC376 [7] were tested for in cellulo M\textsuperscript{Pro} inhibition potency. Nirmatrelvir has a determined in cellulo EC\textsubscript{50} value as 3.4 μM [38], and GC376 shows a in cellulo EC\textsubscript{50} value of >2 μM

Fig. 4. Cellular potency of MPI11-28 in their inhibition of M\textsuperscript{Pro} to drive host 293T cell survival and overall M\textsuperscript{Pro}-eGFP expression.
2.5. Characterisations of antiviral potency of six selected inhibitors on three SARS-CoV-2 variants

Four most cellularly potent inhibitors MPI16, MPI17, MPI125, and MPI26 that contain a P3 O-tet-butyl-threonine and two mildly cellularly potent inhibitors MPI18 and MPI27 that don’t contain a P3 O-tet-butyl-threonine were analyzed on their antiviral potency against three SARS-CoV-2 variants including USA-WA1/2020, Beta and Delta (Fig. 5). We conducted plaque reduction neutralization tests in Vero E6 cells for all inhibitors. We infected Vero E6 cells by virus in the presence of an inhibitor at various concentrations for three days and then quantified viral plaque reduction. As presented in the attached figure and table, all four inhibitors that contain a P3 O-tet-butyl-threonine showed high antiviral potency with determined EC_{50} values around 1 μM and MPI18 and MPI27 that don’t have a P3 O-tet-butyl-threonine showed only moderate antiviral potency. It should be noted that MPI5 containing a P3 valine showed mild cellular M^{Pro} inhibition potency but potent antiviral activity [13,31]. Its antiviral activity may not only come from the inhibition of M^{Pro} but also other mechanisms.

2.6. Cytotoxicity of MPI8, MPI16, MPI17, MPI18, MPI25, MPI26 and MPI27

We characterized cytotoxicity for these seven compounds in 293T cells using the MTT assay [39]. Cytotoxicity curves of these inhibitors are shown in Supplementary Fig. S2. Their determined CC_{50} values are 109.2, >200, >200, 130, 182.9 and 114.0 μM for MPI8, MPI16, MPI17, MPI18, MPI25, MPI26 and MPI27, respectively (Table 1). In term of selectivity index (CC_{50}/antiviral EC_{50}), MPI8, MPI16, MPI17, and MPI18 are around or above 200.

3. Discussion

There are 11 unique proteolytic sites in pp1a and pp1b that are hydrolyzed by M^{Pro}. Glutamine is a strictly required residue at the P1 site for all 11 sites. Due to this strict requirement and medicinal chemistry information learnt from the same function enzyme in SARS-CoV, most peptidyl inhibitors that have been developed for M^{Pro} have maintained a β-(S-2-oxopyrrolidin-3-yl)-alanine analog at the P1 site for improved potency. In our design, we keep opal at this site for all inhibitors. Our crystallography results show a well-shaped opal side chain in all M^{Pro}-inhibitor complexes that fits neatly in the P1 binding pocket of the enzyme. Since opal is optimized for this site, there is small chemical space to manipulate its side chain for improved binding. In all determined structures, the side chain of N142 adopts different conformations that do not necessarily interact with the opal oxopyrrolidine ring. It may be possible to modify the opal oxopyrrolidine ring to introduce interactions such as hydrogen bond(s) with the N142 side chain for improved binding. Among all 11 M^{Pro}-targeted proteolytic sites, 9 have a P2 leucine, 1 has a P2 phenylalanine, and 1 has a P2 valine. The enzyme has an apparent preference for a P2 leucine in its substrates.

Comparison of in vitro enzyme inhibition potency for all MPIs that we have developed also reveals that M^{Pro} prefers a P2 residue with a similar size as leucine in a peptidyl inhibitor for favorable binding. Converting the isopropyl group at the β-carbon of the P2 residue to cyclopropyl and 2-furanyl improves the enzyme inhibition kinetics. Although changing the P2 leucine in an inhibitor to a larger residue such as phenylalanine and β-cyclohexylalanine diminishes in vitro M^{Pro} inhibition potency, the level is not significant. It is interesting to see that a β-isoprenyl group at the P2 residue has no fixed conformation but a cyclopropyl group at the same position has a defined conformation in crystal structures. Since inhibitors with a β-cyclopropyl group at the P2 residue have typically high in vitro enzyme inhibition potency, it is possible that adding an additional group such as methyl to the β-carbon of a P2 β-cyclopropylalanine will lead to better inhibitors. In the crystal structure of M^{Pro}, MPI14, the P2 side chain of MPI14 forms a π-π stacking interaction. This π-π stacking interaction likely contributes to the high in vitro enzymatic inhibition potency of MPI14. Although MPI15 is not soluble, it is expected to have high in vitro enzymatic inhibition potency as well due to a similar π-π stacking interaction that can be formed between MPI15 and M^{Pro}. Other five-membered rings that contain one or more heteroatoms at the P2 β-carbon may also lead to high in vitro enzymatic inhibition potency since a similar π-π stacking interaction can be formed between them and M^{Pro}. In most of our determined M^{Pro}-inhibitor complexes, we noticed that the M49-containing loop of M^{Pro} has weak electron density, indicating that this loop adopts a flexible structure. Since this loop caps the M^{Pro} P2 binding pocket, this flexibility will potentially allow peptide inhibitors with a P2 residue larger than all moieties that have been tested in our inhibitors at this site to be tolerated by M^{Pro}. This probability needs to be explored.

M^{Pro} shows almost no preference toward the P3 residue in its targeted proteolytic sites in pp1a and pp1b. We explored a variety of chemical variants at this site including dialkyl glycines and 1-amino-cyclopropane-1-carboxylate that have no α-proton. Although M^{Pro} displays a clear preference for a P3 valine, other substituents at this site are well tolerated. For most inhibitors, their P3 residues can be well refined. Given the diverse structures of tolerable chemical compositions at this site, α-amino acids might be introduced at this site for the development of novel M^{Pro} inhibitors as well. This will need to be explored further. One interesting observation is that α,α-disubstituted glycines at the P3 position push their adjacent N-terminal carbamate carbonyl group to adopt an orientation rotating about 90° away from a position that has been observed in M^{Pro}-inhibitor structures into the M^{Pro} active site. It will be interesting to see whether this observation can be applied to optimize existing M^{Pro} inhibitors including nirmatrelvir to achieve...
better antiviral potency. All our crystal structures show weak electron density around the N-terminal group indicating its weak binding to the enzyme. Adding an m-chloride to the N-terminal CBZ group also leads to significant decrease in in vitro enzyme inhibition potency. So far, our studies indicate that both CBZ and indole-2-carboxyl are not optimal chemical groups for interactions with Mpro. Some smaller groups need to be explored at this site. One example will be trifluoroacetamide as seen in nirmatrelvir.

Although most inhibitors we have developed in this study have high in vitro enzyme inhibition potency and two inhibitors have IC_{50} values reaching the characterization limit, just a few inhibitors show cellular potency to inhibit Mpro in 293T cells. Inhibitors that have a P3 O tert-butyl-threonine all show strong cellular potency. All other P3 substituents lead to relatively weaker cellular potency, although MPI5, MPI18, and MPI27 showed mild cellular potency. Many factors may contribute to this phenomenon. Since all MPIs are peptidyl inhibitors, a small or native P3 residue might be prone to proteolytic degradation by host proteases leading to low cellular potency. This is partially supported by characterizable cellular potency for MPI18-19 that have a P3 β tert-butyl-alanine. It is also likely that a P3 O tert-butyl-threonine introduces more favorable cellular permeability than other P3 substituents into an Mpro inhibitor. This aspect needs to be further investigated.

MPI21-23 that have a P3 dialkylglycine are expected to be more resistant to proteolytic digestion by human proteases than other MPIs. Their low cellular potency might be due to low cellular permeability. Based on cellular potency of MPIs, we can also derive that an N-terminal CBZ group works better than the other two groups that have been tested at this site and a P2 β-cyclohexylalanine favors high cellular potency as well.

In terms of antiviral potency, results for six tested compounds agree well with their cellular potency, supporting the notion that the inhibition of Mpro in infected cells is enough to achieve the prevention of viral replication. It should be noted that 293T cells were used in the cellular assay and Vero E6 cells were used in the antiviral assay. Different cell lines may have different enzymes which may contribute to different metabolism of inhibitors, leading to the discrepancy between cellular Mpro inhibition potency and antiviral potency. Six tested inhibitors also exhibited relatively low cytotoxicity. With this level of selectivity, potential achievement of their use in containing SARS-CoV-2 in COVID-19 solve both metabolic and cytotoxicity concerns. Therefore, new antiviral strategies for the treatment of COVID-19 might be explored at this site. One example will be trifluoroacetamide as seen in nirmatrelvir.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ejmech.2022.114570.

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