Coronavirus helicases: attractive and unique targets of antiviral drug-development and therapeutic patents

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ABSTRACT

Introduction: Coronaviruses encode a helicase that is essential for viral replication and represents an excellent antiviral target. However, only a few coronavirus helicase inhibitors have been patented. These patents include drug-like compound SSYA10-001, aryl diketo acids (ADK), and dihydroxycromones. Additionally, adamantane-derived bananins, natural flavonoids, one acrylamide derivative ([E]-3-(furan-2-yl)-N-(4-sulfamoylphenyl)acrylamide), a purine derivative (7-ethyl-8-mercapto-3-methyl-3,7-dihydro-1 H-purine-2,6-dione), and a few bismuth complexes. The IC50 of patented inhibitors ranges between 0.82 μM and 8.95 μM, depending upon the assays used. Considering the urgency of clinical interventions against Coronavirus Disease-19 (COVID-19), it is important to consider developing antiviral portfolios consisting of small molecules.

Areas covered: This review examines coronavirus helicases as antiviral targets, and the potential of previously patented and experimental compounds to inhibit the Severe Acute Respiratory Syndrome Coronavirus-2 (SARS-CoV-2) helicase.

Expert opinion: Small molecule coronavirus helicase inhibitors represent attractive pharmacological modalities for the treatment of coronaviruses such as SARS-CoV and SARS-CoV-2. Rightfully so, the current emphasis is focused upon the development of vaccines. However, vaccines may not work for everyone and broad-based adoption of vaccinations is an increasingly challenging societal endeavor. Therefore, it is important to develop additional pharmacological antivirals against the highly conserved coronavirus helicases to broadly protect against this and subsequent coronavirus epidemics.

1. Introduction

Helicases are ubiquitous nucleic acid unwinding enzymes. These biological motors couple the chemical energy of nucleotide triphosphate hydrolysis (NTPase) to mechanical energy that translocates through nucleic acids, unwinding the helical structure as it progresses, thus the term ‘helicase.’ Efficient genome replication, recombination and repair require single stranded DNA (ssDNA) or single stranded RNA (ssRNA) as a template that is largely devoid of secondary structures [1]. Helicases in situ generate ssDNA or ssRNA, and due to this crucial role during genome replication, repair and recombination, defects in helicase function can lead to many genetic disorders. Notable examples of helicase-associated disorders include Bloom’s syndrome, Werner’s syndrome, and X-chromosome-linked α-thalassemia [2–9].

Similar to their hosts, viruses rely on nucleic acid unwinding for efficient replication of their genetic material. While many viruses encode helicases, notable exceptions such as HIV-1 do not encode helicases, relying upon host cell helicases to facilitate their replication. Some viruses encode helicase as full-length individual proteins, whereas others encode helicase domains within other functional proteins. All identified CoVs encode a full-length helicase [10–13], whereas other highly pathogenic viruses such as flaviviruses (hepatitis C, dengue and Zika viruses) encode a singular multifunctional protein encoding protease and helicase activities within a single mature protein [14,15].

To date, seven human coronaviruses (hCoVs) have been discovered. These are hCoV-229E, NL63, OC43, HKU1, Severe Acute Respiratory Syndrome coronavirus (SARS-CoV), Middle East Respiratory Syndrome coronavirus (MERS-CoV), and most recently SARS-CoV-2. Of these, hCoV-229E, NL63, OC43, and...
HKU1 cause mild self-limiting respiratory infections, whereas SARS-CoV, MERS-CoV and SARS-CoV-2 cause highly infectious and deadly diseases.

CoVs are large (15–30 kb) positive (+) single-stranded RNA viruses. The polyadenylated genome has open reading frames (ORFs) that encode pp1a and pp1ab through a – 1 ribosomal frameshift during translation [16,17]. These large polyproteins are proteolytically cleaved to generate mature CoV non-structural proteins (nsps) as well as the structural and accessory proteins [10,18–24]. The polyproteins are processed by a virus-encoded papain-like proteinases (PLpro; within nsp3) [25] and nsp5 (3 CLpro) [10,16,19,26–29] to yield up to 16 nsps with diverse functions and up to 10 structural proteins that facilitate viral assembly [30–34]. The singular helicase protein of hCoVs is encoded by nsp13 [12]. nsp13, in concert with additional nsp proteins, assembles into a replication-transcription complex that binds to the 3’ untranslated region of the genomic RNA and generates (-) strand RNA as well as subgenomic RNAs [30–34]. A recent review has covered patented inhibitors of different CoV targets [35], here we have focused on the patented inhibitors of nsp13.

2. SARS-CoV-2 life cycle and steps of nsp13 inhibition

SARS-CoV-2 infection initiates with the attachment of the virus through the cell surface receptor ACE2 interacting with the receptor binding domain (RBD) of Spike protein (S-protein). The host protease TMPRSS2 acts synergistically during the initial step of viral attachment to the cell [36] as it cleaves the viral S-protein into S1 and S2, triggering endocytosis of the viral particle (Figure 1). The pH of the cytoplasm induces endosomal acidification and subsequent release of the viral genome into the cytoplasm. The host ribosomal machinery translates the (+) sense viral RNA genome into polyproteins pp1ab and pp1b. The autocleavage of viral polyproteases, M\textsuperscript{pro} and 3C\textsuperscript{pro}, is followed by the cleavage of pp1ab and pp1b polyproteins into ~16 non-structural proteins (nsps) and ~10 structural proteins. The replicative nsps form a complex that copies the (+) sense viral RNA genome into the (-) sense RNA genome. The nascent (-) sense RNA is then used by a replicase complex comprised of nsps within the double membrane vesicles (DMVs), and generates the (+) sense genomic and subgenomic RNAs. The subgenomic RNAs are translated into the structural proteins that assemble at the Endoplasmic Reticulum Golgi Intermediate Compartment (ERGIC). The N-protein coated RNA genome and structural protein containing cell membrane form the virus like particle (VLP), which egresses from the cell as a mature and infectious virion.

The published reports have shown that nsp13 is essential for replication of the viral genome [12]. nsp13 also interacts with other nsps including nsp7, nsp8, nsp9, nsp12, and nsp14 [37–42]. Recent structural studies have confirmed that nsp13 form complex with nsp7, nsp8, and nsp12 as well as with nsp7, nsp8, nsp9, and nsp12 [36,43]. Additionally, recent data show that SARS-CoV-2 nsp13 interacts with host factors involving in different signaling pathways [44–46]. A definitive role of nsp13 during (+) sense RNA or (-) sense RNA or both steps is still not established. However, SSYA10–001, a patented nsp13 inhibitor, inhibited (+) sense, but not (-) sense, RNA synthesis in a SARS-CoV replicon assays, suggesting that nsp13 is required for the (+) strand synthesis. While it is clear that nsp13 is a pleiotropic factor, more data are needed to establish a definitive role of nsp13 in viral RNA synthesis.

3. Structure-function relationship of CoV helicases

Using conserved sequence motifs, helicases have been classified into six superfamilies (SF1–SF6) [12,47,48]. The nsp13 of hCoV and Mouse Hepatitis Virus (MHV) (a well-studied CoV) belong to the SF1 superfamily [12,48,49]. While, the function of nsp13 CoV during viral replication is not well understood, studies have shown that it is essential for viral replication (see below). Biochemical and molecular biological studies have shown that nsp13 interacts with nsp7, nsp8 and nsp12 [37–40], and a recent cryoEM structure showed that SARS-CoV-2 nsp13 associates with nsp12 when complexed with nsp8 [49]. nsp8 appears to function as an interaction hub for SARS-CoV nsp8 [39]. Nsp13 is a multifunctional protein that has 5’-3’ directed helicase [12,49–49–51], ATPase and RNA 5’-triphosphatase activities [52].

The crystal structure of MERS-CoV nsp13 was the first full-length high-resolution structure reported for any hCoV [53]. Subsequent crystal structure of SARS-CoV nsp13 [54] showed that the nsp13s from the two viruses share a high degree of structural similarity. These structures also revealed that nsp13 consists of multiple domains including: a zinc binding domain (ZBD); and an SF1 helicase core connected by the ‘Stalk’ and 1B domains (Figure 2). The catalytic functions of nsp13 are conducted by the SF1 helicase core formed by two RecA-like domains (RacA1 and RecA2) (Figure 2). The N-terminal ZBD is also essential for nsp13 function [55] as several ZBD mutations inactivate ATPase and/or helicase activities of equine arteritis virus (EAV) nsp10 (the functional equivalent to SARS-CoV-2 nsp13). Inactivation of ZBD blocks sgRNA (segmented RNA) and genomic RNA synthesis as well as the production of infectious progeny [55,56]. These results suggest that the ZBD is required for either the structural integrity of nsp13 or for the efficient replication of the virus or both.

The structures of SARS-CoV, MERS-CoV and SARS-CoV-2 nsp13 not only provide valuable insights into nsp13 macromolecular organization, but also pave the way for identifying potential.
small molecule inhibitor pockets (Figure 3). Examples of these pockets, computed by the SiteID program of SYBYL (version X, Certara, St. Louis, MO), are shown as yellow space-filled atoms (Figure 3). Majority of binding pockets are in the close vicinity of the highly conserved helicase motifs (Figure 3). This is not surprising since these motifs are supposed to bind ATP and/or RNA substrate. The structure was solved in the absence of substrates, it is possible that the binding sites of these substrates remained unoccupied. Additionally, potential small molecule binding pockets were detected close to K508 and Y277. Importantly, mutations K508A and Y277A conferred resistance to SSYA10–001.

4. CoV helicases as antiviral targets

CoV nsp13 is a highly conserved protein as there is only a single amino acid difference between nsp13s of SARS-CoV (I570) and SRS-CoV-2 (V570). A phylogenetic tree derived from the sequence alignment of nsp13s of seven hCoVs, one bat (RaTG13) and MHV (Figure 4) shows that (i) SARS-CoV-2 is closely related to bat CoV RaTG13 (purple shaded node) as reported previously [58–60], and (ii) all six SF1 helicase motifs are highly conserved. A structural analysis showed that all six conserved motifs are sequestered at the interface of RecA1
and RecA2 domains and form nucleotide/RNA binding sites [53]. Additionally, many potential small molecule binding sites are near RecA1/RecA2 interface (Figure 3), and within the vicinity of conserved motifs (Figure 3), it can be easily postulated that (i) CoV nsp13 are attractive antiviral targets and (ii) the compounds binding in the pockets near conserved motifs may serve as pan-coronavirus inhibitors.

5. Patented nsp13 inhibitors

5.1. SSYA10–001

One of the most characterized CoV nsp13 patented inhibitor is SSYA10–001 (US20140005241A1; 2016). SSYA10–001 is a 1,2,4-triazole compound (Figure 5) that was identified through a screening of a library of drug-like compounds using fluorescence resonance energy transfer (FRET) based helicase assay [63]. SSYA10–001 functionally inhibited helicase and ATPase activities of SARS-CoV nsp13 in a noncompetitive manner with respect to ATP and nucleic acid substrate [63]. Subsequent cell-culture assays showed that SSYA10–001 inhibits SARS-CoV, MERS-CoV and MHV with comparable IC₅₀ [64]. Therefore, it was termed as a pan-CoV inhibitor [64]. To identify compounds that share structural and physico-chemical properties of SSYA10–001, we conducted a PubChem search using SSTA10–001 as template. This search resulted in the identification of seven compounds (Table 1). Below we discuss structural and physico-chemical properties of these compounds with respect to SSYA10–001, however, they are yet to be tested for anti-CoV activity.

**Figure 2. Structure of SARS-CoV nsp13.** This figure shows the overall structure and domain organization of SARS-CoV nsp13. This figure was generated by PyMol [57].

**Figure 3.** Panel A shows potential small molecule inhibitor binding pockets (yellow space-filled atoms) in the structure of SARS-CoV as determined by SiteID program (Tripos Associates, St. Louis, MO). These binding sites were calculated from the crystal structure of SARS-CoV nsp13 (PDB file 6JYT [47]) after changing selenomethionine residues to methionine. Panel B shows the close up of conserved motifs, computed small molecule binding pockets and SSYA10–001 binding site residues K508 and Y277 (shown as ball-and-sticks). The dotted line represents (42 Å) the distance between two Cα-atoms of Y277 and K508. Residues that interact with K508 from Motifs IV and V are also shown in this panel. The carbon, nitrogen and oxygen atoms are colored as white, blue and red, respectively. This figure was generated by Sybyl X (Certara, St. Louis, MO).
Compounds
th CoVs, 13,953,241
4-Ethyl-5-(2-phenylethyl)-4-3-
(2-Nitrophenyl)sulfanylmethyl]-4-
prop-2-enyl-1-
757,359
CPD-062
7,130,850
CPD-850
3,687,815
CPD-815
900,999
CPD-999
1-
H-1,2,4-triazole-3-thioli
7,130,850
CPD-850
4-Allyl-5-(2-piperidin-1-ylethyl)-4-
H-1,2,4-triazole-3-thiol
675,062
CPD-062
4-Ethyl-5-(2-phenylethyl)-4-
H-1,2,4-triazole-3-thiol
13,953,241
CPD-241
4-Chloro-4'-methoxy-2-nitroazobenzene
757,359
CPD-359
N-Ethyl-2-methyl-N-[(E)-3-(2-nitrophenyl)
prop-2-enyl]prop-2-amine

Figure 4. Phylogenetic analyses of CoVs and conservation of motifs among selected CoVs. The dendrogram shows the evolutionary relationship among hCoVs, MHV and RaTG13 using amino acid sequence of nsp13. Note that SARS-CoV-2 is closely related to RaTG13 (purple highlighted node). This figure also shows the sequence alignment of six conserved nsp13 motifs together with SSYA10–001 binding site residues marked as asterisks (*). The amino acid sequence is color coded. Dash (-) represents the gap in the sequences of nsp13, and AA represents amino acids. The figure was generated using the R package ggtree [61]. The R programming language [62] was used to generate this figure, Figures 6 and 7. The in-house generated R codes for these figures are available upon request.

Figure 5. Structure of SSYA10–001 and related compounds. The compounds were searched from PubChem database using SSYA10–001 as template. The compounds were scored using Tanimoto fingerprints [66].

Table 1. List of compounds in PubChem sharing structural similarity with SSYA10–001.

| Compound | PubChemCID | Chemical Name |
|----------|------------|---------------|
| SSYA10–001 | 2,807,230 | 3-{[2-Nitrophenyl)sulfanylmethyl]-4-prop-2-enyl-1-H-1,2,4-triazole-5-thione |
| CPD-673 | 86,246,673 | 1-{[2-Isocyanophenyl)sulfanylmethyl]-2-nitrobenzene |
| CPD-815 | 3,687,815 | 4-Allyl-5-(2-phenylethyl)-4-H-1,2,4-triazole-3-thiol |
| CPD-999 | 900,999 | 5-{[2-Nitrobenzyl)sulfanyl]-1-methyl-1-H-tetrazole |
| CPD-850 | 7,130,850 | 4-Allyl-5-(2-piperidin-1-ylethyl)-4-H-1,2,4-triazole-3-thiol |
| CPD-062 | 675,062 | 4-Ethyl-5-(2-phenylethyl)-4-H-1,2,4-triazole-3-thiol |
| CPD-241 | 13,953,241 | 4-Chloro-4'-methoxy-2-nitroazobenzene |
| CPD-359 | 757,359 | N-Ethyl-2-methyl-N-[(E)-3-(2-nitrophenyl)prop-2-enyl]prop-2-amine |

Clustering of compounds by structural or property similarity is a powerful approach for correlating compound features with biological activities. This technique also provides a glimpse of structure activity relationship when combined with binding/inhibition of the target protein. A heatmap derived from the structure-based hierarchical clustering of compounds in Table 1 generated by ChemMine Tools [65] is shown in Figure 6. To generate structure-based clustering of these compounds, the atom pair descriptors (features) for each compound are generated and used to calculate a similarity matrix based on the common and unique features observed among all compound pairs using the Tanimoto coefficient [66]. The Tanimoto coefficient ranges between 0 and 1 with a higher value representing greater similarity. In subsequent clustering steps, the similarity matrix is converted into a distance matrix (Z-Score) by subtracting the Tanimoto similarity values from 1. The Z-scores were finally used to clustering the compounds (Figure 6). From this heatmap, it is clear that compound CPD-850 (Figure 5) is most dissimilar to the other compounds, while CPD-815 and CPD-062 are very similar with respect to structural properties, and a high degree of similarity exists between SSYA10–001, CPD-815 and CPD-062 (Figures 5 and 6).

A heatmap generated using physicochemical property-based clustering of compounds in Table 1 is shown in Figure 7A.
Selected Open Babel descriptors [67] (HBD, number of H-bond donors; HBA, number of H-bond acceptors; ArBD, aromatic bonds; and MW, molecular weight) were used to generate the heatmap and dendrogram using the ChemMineR package (Cheminformatics Toolkit for R) [68] (Figure 7A). This heatmap highlights the correlations between the compounds in Table 1. For example, ‘ArBD’ (aromatic bond) feature reveals that 6 out of 8 compounds have comparable aromatic bonds, a measure of hydrophobicity. Additionally, the computed MW and logP of compounds is closely related. The dendrogram at the top of the figure shows that the hydrophobicity of the compounds is closely related to the molecular weight of the compounds. Additionally, horizontal dendrograms show that the identified compounds can be grouped, and prioritized for experimental evaluation. For example, CPD-241 is closely related to SSYA10–001, although CPD-241 is significantly hydrophobic (as seen from logP) compared to SSYA10–001. To get better insight about the solubility of all compounds compared to SSYA10–
001, the plotted logP values are shown in Figure 7B. It is clear that except CPD-241, each compound has a better or comparable solubility profile compared to SSYA10–001. Additionally, the docking of these compounds at the two possible binding sites in the structure of nsp13 coupled with the experimental data can provide SAR strategies to develop more potent inhibitors.

5.2. Dihydroxychromone and aryl diketoacids

Two Korean patents include compounds based on 3, 5-dihydroxychromone and aryl diketoacids (Figures 8 and 9) (KR20100029528A; 2010 and KR20110006083A; 2011). These patents arose from the identification of natural flavonoids myricetin, scutellarein and quercetin as potent inhibitors of SARS-CoV nsp13 [69,70] (Figure 8). The compounds described in these patents contain common scaffold 3, 5-dihydroxychromone. Using this scaffold, modified quercetin derivatives were developed that showed anti-nsp13 activity. Modifications \( R_1 = R_2 = H \) and \( R_3 = 3\text{-CN-Benzyl} \), and \( R_3 = 4\text{-Cl-Benzyl} \) were the most potent nsp13 inhibitors with IC\(_{50} \sim 3.0 \) µM.

The diketoacids are known to chelate two divalent cations, and thereby inhibit enzymes that use metal ions (such as Mg\(^{2+}\) and Mn\(^{2+}\)) as cofactors. For example, all approved HIV-1 integrase inhibitors are diketoacid compounds (e.g., Dolutegravir, Figure 9) [71]. The inventors of Korean patent (KR20100029528A; 2010) exploited the divalent cation chelating property of diketoacids and developed a series of compounds that inhibited nsp13 in in vitro assays. The most potent inhibitor was a compound with \( R_1 = H, R_2 = \text{NHCH}_2(4\text{-Cl-phenol}) \), and \( R_3 = H \) (1-ADK, Figure 9). This compound had an IC\(_{50} \) = 0.96 µM, and EC\(_{50} \) = 0.82 µM against SARS-CoV.

5.3. Mechanism of SSYA10–001, dihydroxychromone and aryl diketoacids inhibition of nsp13

The SSYA10–001 mechanism of nsp13 inhibition, the functional impact of mutations within the predicted SSYA10–001 binding site was examined. Two nsp13 mutations, Y277A and K508A, conferred resistance to SSYA10–001, as their estimated IC\(_{50S} \) for helicase function were 12 and 50 µM, respectively.
compared to 5.9 μM for WT SARS-CoV nsp13 [64]. Based on these results and the computational modeling, it was concluded that Y277 and K508 were part of the binding pocket for SSYA10–001 within SARS-CoV nsp13 [64]. However, the crystal structure of SARS nsp13 showed that the two residues (Y277 and K508) are nearly 42 Å apart (Figure 3), suggesting that the two residues are not part of the same binding pocket. Hence, the precise SSYA10–001 binding pocket remains unknown. It is possible that SYA10–001 (i) binds at one site and exerts a long-range effect on the other site or (ii) there exist two binding sites for SSYA10–001.

In the crystal structure of SARS-CoV nsp13 (PDB file 6JYT [54]), K508 is exposed to the solvent and does not appear to contribute to the nearest small-molecule binding pocket. However, K508 forms a salt-bridge with D542 (motif IV), which in turn forms a salt-bridge with K569 (motif V) (Figure 3). Mutation K508A may disrupt the positions of D542 and K569. Importantly, D542 is next to Y543, whereas K569 is next to I570. Both residues (Y543 and I570) hydrophobically interact with W506, which is part of the proposed SSYA10–001 binding pocket [64]. Hence, it appears that the resistance to SSYA10–001 due to K508A mutation is an indirect effect structural alteration in its vicinity. Mutation Y277A is resistant to SSYA10–001 although ~twofold less compared to K508A. The SitelD program of Sybyl X (Certara, St. Louis, MO) predicted a small pocket near Y277. However, the size of this pocket appeared too small to bind SSYA10–001. At this point, the structure nsp13 does not provide a plausible mechanism for SSYA10–001 resistance of Y277A. A crystal structure of nsp13/SSYA10–001 is needed to resolve this issue. The molecular mechanism of inhibition of other patented nsp13 inhibitors (aryl diketoacid and 3,5-dihydroxycromone) has not been investigated and the binding sites for these inhibitors are currently unknown. However, based on the metal binding properties of these compounds, it is quite possible that these compounds bind at or near motifs I and II.

### 6. Miscellaneous nsp13 inhibitors

In addition to above mentioned patented nsp13 inhibitors, several compounds that inhibited CoV nsp13 activities have been reported [69,70,72–80]. These include adamantane-derived bananins, bismuth complexes, thioxopyrimidine derivatives, an acrylamide derivative (E)-3-(furan-2-yl)-N-(4-sulfamoylphenyl) acrylamide, a purine derivative (7-ethyl-8-mercaptopo-3-methyl-3,7-dihydro-1 H-purine-2,6-dione), and RNA aptamers [81]. Of all tested adamantane-derived bananins, bananin was the most potent with an IC$_{50}$ of 2.3 μM and 3.0 μM for ATPase and helicase activities of SARS-CoV nsp13, respectively [66]. An acrylamide derivative (E)-3-(furan-2-yl)-N-(4-sulfamoylphenyl)acrylamide showed an IC$_{50}$ for ATPase activity comparable to bananin, but the IC$_{50}$ for helicase activity was 4.4-fold greater (13.0 μM) [72]. The inhibitory profile of a purine derivative (7-ethyl-8-mercaptopo-3-methyl-3,7-dihydro-1 H-purine-2,6-dione) was poorer than bananin (ATPase IC$_{50}$ = 8.66 ± 0.26 μM and helicase IC$_{50}$ = 41.6 ± 2.3 μM) [73]. Some of the bismuth complexes had IC$_{50}$ values for ATPase activity of SARS-CoV nsp13 less than 1 μM and IC$_{50}$ values for helicase activity less than 3 μM [75]. However, these compounds have not been pursued since their discovery in 2007 until very recently it was shown that they inhibit SARS-CoV-2 nsp13 [82]. Very recently, another compound N-(4-(4,6-dioxo-2-thioxotetrahydroprymidin-5(2 H)-ylidene)ethyl)phenyl)acetamide showed good potency for the inhibition of ATPase activity of SARS-CoV nsp13 (IC$_{50}$ = 1.19 ± 0.16 μM) [80]. However, the IC$_{50}$ (32.9 ± 1.0 μM) for helicase function was similar to that of 7-ethyl-8-mercaptopo-3-methyl-3,7-dihydro-1 H-purine-2,6-dione [80].

### 7. Expert opinion

Here, we present a description and analysis of the currently patented inhibitors of nsp13 as potential small molecule inhibitors of SARS-CoV or MERS-CoV. Considering the high sequence homology between nsp13 of SARS-CoV-2 and SARS-CoV (98.9%), we suggest that it is extremely likely that the compounds discussed in the review will inhibit SARS-CoV-2 nsp13. Mechanistically, among the patented molecules, SSYA10–001 is the best characterized nsp13 inhibitor. The existing analogs of SSYA10–001 have not been tested for efficacy against SARS-CoV or SARS-CoV-2 nsp13 inhibition, however, since they are structurally related and may bind the same nsp13 region, it is possible that one or more analogs may show better efficacy than SSYA10–001.
Despite the critical role of helicases, drug-discovery against this class of enzymes has lagged behind when compared to other viral targets, including nucleic acid polymerases. However, the clinical success of herpes simplex virus (HSV) helicase/primase inhibitors Amenamevir (ASP2151) and Pritelivir (AIC316) demonstrates the potential of helicases as small molecule antiviral targets [83–85]. Following the encouraging results of clinical trials of Pritelivir [86], the Food and Drug Administration (FDA) approved this compound for the treatment of acyclovir-resistant mucocutaneous herpes simplex virus (HSV) infections in immunocompromised adults [87]. Efforts have been made to identify the inhibitors of helicase from other viruses as reported in prior reviews [88,89]. For example, the hepatitis C virus NS3 helicase is an attractive target for small molecule antiviral compounds [90] as several lead compounds have been identified. One class of compounds, triazoles, inhibit HCV helicase activity and impair cellular HCV RNA replication and appear to hold immense potential. Unfortunately, these compounds failed to enter the clinic.

Due to their essential function(s) during the viral life cycle, helicases are clearly attractive antiviral targets regardless of the pathogen. While some helicase-targeting small molecules inhibitors have entered clinical trials, drug development against helicases remains challenging. One of these challenges is to design specific nsp13 inhibitors that compete with the natural substrate (ATP) and bind at the ATP binding site. As with many viral functional proteins and enzymes, the ATPase function of nsp13 is common to a broad array of cellular enzymes. Therefore, not only is binding essential, but substrate specificity amongst a sea of similar substrates adds an additional layer of complexity. For example, protein phosphatase 2A (PP2A), which is involved in many cellular functions, also conducts ATP hydrolysis by binding ATP through two metal ions [91]. Hence, any inhibitor that chelates metal ions can bind PP2A-like enzymes resulting into adverse toxicity profiles. Additionally, a two metal ions mechanism originally proposed for the 3′ – 5′ exonuclease function of E. coli DNA polymerase I [92] is used by many cellular polymerases and primases. Therefore, the compounds targeting ATPase function of nsp13 through metal chelating property can potentially bind to these enzymes and interfere normal cellular function. This type of challenge is not unique to drugs targeting viral proteins/enzymes, as similar issues emerged during the discovery of anti-HIV compounds. Several nucleoside reverse transcriptase inhibitors (NRTIs) inhibit polymerase γ, a mitochondrial DNA polymerase and exert mitochondrial toxicity [93–96].

The structures of viral components such as nucleic acid polymerases, proteases and integrases have helped overcome the specificity challenges. For example, prior to the structure of HIV-1 integrase being solved, the structures of Prototype foamy virus integrase complexed with DNA was used as a molecular scaffold to identify potent integrase inhibitors [96–98].

Currently patented nsp13 inhibitors have limitations. First, the potency of these compounds is in the micromolar range. Second, the pharmacokinetics data for all patented compounds are not known. And most importantly, ADK (aryl diketo acid) and dihydroxychromones contain chemical moieties that can inhibit the function of many cellular enzymes as they can potentially scavenge metal ions required for many cellular enzymes. These limitations can be overcome by the combination of structure-based drug design techniques, medicinal chemistry, and virological assays. Thus, the reported structures of SARS-CoV [54] and MERS-CoV [53] and recently reported structure of SARS-CoV-2 in complex with nsp12, nsp7 and nsp8 [49] as well as the structures of helicases from other CoVs [99,100] provide important opportunities to design specific CoV inhibitors. The resistance profile, most likely obtained through the virus passage assays can aid in the identification of inhibitor binding site residues that can be used to exploit the structures of nsp13. Such strategies have been previously used to design non-nucleoside reverse transcriptase inhibitors (NNRTIs) and the mechanism of their resistance [101–105]. Additional challenges in designing nsp13 inhibitors is the constant evolution of SARS-CoV-2 virus. The diversity in viral genome is known to compromise the efficacy of antiviral drugs [106,107]. Continuously evolving SARS-CoV-2 including two mutations (P504L and Y541C) [108] can also limit the efficacy patented and investigational nsp13 inhibitors.

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