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Functional and druggability analysis of the SARS-CoV-2 proteome

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Abstract

The infectious coronavirus disease (COVID-19) pandemic, caused by the coronavirus SARS-CoV-2, appeared in December 2019 in Wuhan, China, and has spread worldwide. As of today, more than 46 million people have been infected and over 1.2 million fatalities. With the purpose of contributing to the development of effective therapeutics, we performed an in silico determination of binding hot-spots and an assessment of their druggability within the complete SARS-CoV-2 proteome. All structural, non-structural, and accessory proteins have been studied, and whenever experimental structural data of SARS-CoV-2 proteins were not available, homology models were built based on solved SARS-CoV structures. Several potential allosteric or protein-protein interaction druggable sites on different viral targets were identified, knowledge that could be used to expand current drug discovery endeavors beyond the currently explored cysteine proteases and the polymerase complex. It is our hope that this study will support the efforts of the scientific community both in understanding the molecular determinants of this disease and in widening the repertoire of viral targets in the quest for repurposed or novel drugs against COVID-19.

1. Introduction

In the past decades, two highly pathogenic coronaviruses (CoVs) known as the severe acute respiratory syndrome coronavirus (SARS-CoV), and the Middle East respiratory syndrome coronavirus (MERS-CoV), triggered global epidemics in 2003 and 2012, respectively (Paules et al., 2020). A new CoV infectious disease (COVID-19), caused by the new pathogenic SARS-CoV-2 appeared in December 2019 in Wuhan, China, spread rapidly worldwide, and was declared a pandemic on March 11th, 2020 by the World Health Organization (WHO). As this work is being written, there have been over 46 million infected cases around the world, with more than 1.2 million fatalities.

Although SARS-CoV-2, SARS-CoV, and MERS-CoV belong to the same genus, SARS-CoV-2 seems to be associated with milder infections. Their fatality rates were 2.3%, 9.5%, and 34.4%, and their basic reproductive numbers (R0) 2.0–2.5, 1.7–1.9, and <1, respectively (Petrosillo et al., 2020). Like SARS and MERS, the typical clinical presentation of severe cases of COVID-19 is pneumonia with fever, cough, and dyspnea. The vast majority of COVID-19 patients have mild disease (around 80%), a lesser percentage (~15%) have moderate disease (with dyspnea, hypoxia or pneumonia with over 50% lung parenchyma involvement), and even a smaller proportion (~5%) of patients develop severe disease with respiratory failure, shock or multiorgan failure (Wu et al., 2020a).

Although the optimal strategy to control this disease would be through vaccination capable of generating a long-lasting and protective immune response, there is an urgent need for the development of a treatment, dictated by: i) the rapid spread of the virus; ii) the increasing number of infected patients with a moderate to severe symptoms that must be treated worldwide; iii) the associated risk of death and depletion of the health systems, particularly in low-income countries.

In terms of antiviral strategies, while no specific therapeutics are available, approved or experimental drugs developed for other diseases are being tested in different clinical trials, in an effort to rapidly find accessible treatments with already established safety profiles, through drug repurposing strategies (Barrantes, 2020; Cavasotto and Di Filippo, 2020).
2020; Villoutreix et al., 2020; Zhou et al., 2020). Most of these strategies have been focused in targeting the two viral proteases, and the RNA-dependent RNA polymerase (RdRp) complex. However, protease inhibitors might bind to host proteases, thus resulting in cell toxicity (Kang et al., 2020), and the effectiveness of nucleoside inhibitors targeting RdRp is decreased by the highly efficient SARS-CoV-2 proofreading machine. Therefore, alternative therapeutic options to fight COVID-19 are urgently needed.

In this context, the characterization of the druggability of the SARS-CoV-2 proteome is of the utmost importance. In this work, we surveyed the functional role of each SARS-CoV-2 protein, analyzed its key structural features, and using either experimental structural data or homology models, we performed a thorough in silico druggability assessment of the viral proteome. Current drug discovery efforts on the proteases and the RdRp complex with in vitro or in vivo experimental validation are also briefly commented, mainly as a complement to our description of targetability. In addition to the catalytic sites of several proteins, we found potential druggable allosteric and protein-protein interaction (PPI) sites throughout the whole proteome.

We do hope our contribution will help in the development of fast and effective SARS-CoV-2-centered therapeutic options, which considering the high similarity among CoVs, might also be effective against related viruses. Moreover, since it is unlikely that SARS-CoV-2 will be the last CoV to threaten global health, those therapeutics might be instrumental in fighting future epidemics.

2. Computational Methods

2.1. Molecular system setup

All structures were downloaded from the Protein Data Bank (PDB) and prepared using the ICM software (MolSoft LLC, San Diego CA, 2019) in a similar fashion as in earlier works (Cavasotto and Aucar, 2020). Succinctly, hydrogen atoms were added, followed by a short local energy minimization in the torsional space; the positions of polar and water hydrogens were determined by optimizing the hydrogen-bonding network, and then all water molecules were deleted. All Asp and Glu residues were assigned a –1 charge, and all Arg and Lys residues were assigned a +1 charge. Histidine tautomers were chosen according to their corresponding hydrogen-bonding pattern.

2.2. Homology modeling

A crude model was built using the backbone structure of the template, and then refined through local energy minimization using ICM (Domenech et al., 2013). To avoid pocket collapse, and taking into account the complete binding site conservation, whenever available, ligands were kept within the binding site during the refinement process, in a ligand-steered modeling fashion (Cavasotto and Palomba, 2015; Phatak et al., 2010).

2.3. Hot-spots and cryptic sites determination

Identification of binding energy hot-spots was performed using FTMap (https://ftmap.bu.edu/) (Kozakov et al., 2015a). The FTMap algorithm samples a library of 16 small organic probe molecules of different sizes, shapes and polarities on the protein using rigid docking. For each probe, all the poses generated are clustered using a 4 Å clustering radius, and then clusters are ranked on the basis of their average energy, keeping the six lowest-energy clusters for each probe. After the probe clusters of the 16 molecules have been generated, they are then re-clustered into hot-spots or consensus sites (CSs) based on vicinity. These CSs are ranked on the basis of the number of their probe clusters. The program offers a protein-protein interaction (PPI) mode, where the aim is to identify hot-spots on protein-protein interfaces. To identify binding sites, the top ranking CS is considered the kernel of the site, and is expanded by adding neighboring CSs with a center-to-center distance (CD) of less than 8 Å to any existing CS, until no further expansion is possible. The binding site is defined as those residues within 4 Å of any probe of the CSs used. The top ranking CS is removed and the procedure repeated starting from the second ranking CS, and so forth. Considering the druggability criteria (see below), only CSs with at least 13 probe clusters were considered to be expanded.

Cryptic sites on proteins were determined using CryptoSite (Cimmermanc et al., 2016) (https://modbase.compbio.ucsf.edu/cryptosite/); cryptic sites are those formed only in ligand-bound structures, but usually “hidden” in unbound structures. Taking into account the analysis of the druggability of cryptic sites, only those cryptic sites with at least 16 probe clusters (determined with FTMap) within 5 Å were considered as potentially druggable (Vajda et al., 2018).

The ICM Pocket Finder method (implemented in the ICM software) predicts the position and shapes of cavities and clefts using a transformation of the Lennard-Jones potential by convolution with a Gaussian kernel of a certain size, and construction of equipotential surfaces along the maps of a binding potential (Abagyan and Kufareva, 2009).

2.4. Druggability criteria

The druggability of a site was characterized based on the CSs generated by FTMap in terms of: i) the number of probe clusters in the primary hot-spot (S), ii) if there are one or more secondary spots with a CD < 8 Å from the primary spot, and iii) the maximum dimension (MaxD) of the connected ensemble (measured as the distance between the two most distant probe atoms within the probe clusters) (Kozakov et al., 2015b). In general, a site was considered druggable if S > 16, CD < 8 Å, and MaxD >10 Å; non-druggable if S < 13 or MaxD < 7 Å; borderline druggable if 13 ≤ S < 16 and CD < 8 Å, or 13 ≤ S < 16, CD ≥ 8 Å, and MaxD ≥ 10 Å.

3. The SARS-CoV-2 viral machinery

The SARS-CoV-2 genetically clusters with the β-coronavirus genus, and is phylogenetically related to SARS-CoV. It is a positive single-strain RNA (+ssRNA), of ~30 kbs (Wu et al., 2020), and it is enclosed in a spherical lipidic bilayer membrane. About the first two-thirds of the viral RNA genome contain the open reading frames (ORFs) 1a and 1ab, which are translated into the polyproteins (pp) 1a and 1ab, which contain the non-structural proteins (nsp’s). The remaining viral genome encode accessory proteins and four essential structural proteins (Cui et al., 2019): the spike (S) receptor binding glycoprotein (ORF2); the nucleocapsid (N) protein (ORF3a); the membrane (M) protein (ORF5), which is a transmembrane (TM) protein involved in the interaction with N; and a small envelope (E) protein (ORF4c), which participates in membrane stability and virus assembly (Si et al., 2008) (Fig. 1). The accessory proteins act co-opting host factors, shutting down host functions to redirect resources to viral replication, avoiding immune responses, and inducing pathogenicity.

Once in a new host, the viral life-cycle consists basically in four stages: virus entry into a host cell, RNA translation and protein processing, RNA replication, and viral particle assembly and release.

During virus entry, the multidomain S protein binds to the host receptor ACE2, it is proteolyzed by host proteases and activated, and thus triggers a series of events that result in the fusion of the viral membrane with the host membrane, and the subsequent release of the viral genome into the host cytoplasm (Cai et al., 2020) (Fig. 1).

The second step in the virus cycle is the translation of viral structural, non-structural, and accessory proteins. Since CoVs particles do not contain any replicase proteins, the translation of viral proteins is the critical step in producing all the necessary machinery for virus RNA replication and assembly. The nsp’s are translated as the large pp pp1a and pp1b, and are cleaved by viral proteases to produce fully active
proteins. CoVs possess the papain-like protease (PL\textsubscript{pro}), and the 3-chymotrypsin-like protease (3CL\textsubscript{pro}) or main protease (M\textsubscript{pro}). The PL\textsubscript{pro} (a domain of nsp3) cleaves nsp1, nsp2 and itself; while M\textsubscript{pro} (nsp5) cleaves the remaining nsp’s, including itself, resulting in total 16 nsp’s (Fig. 2B).

In CoVs the RNA replication takes place in double membrane vesicles (DMVs), colloquially known as viral factories (Snijder et al., 2020). DMVs are membrane structures derived from the Endoplasmic Reticulum (ER), formed through the action of proteins nsp3, nsp4 and nsp6 (Angelini et al., 2013). Due to the ability of some nsp3 domains to bind RNA and to the N protein, the viral RNA and the entire replication machinery are located in the DMVs (Snijder et al., 2020). Furthermore, DMVs constitute a barrier that prevents viral RNA from being detected by the cell’s antiviral response machinery (den Boon and Ahlquist, 2010).

The third step in the viral cycle is the replication of the viral RNA. As

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**Fig. 1.** Schematic representation of a SARS-CoV-2 viral particle and key steps in virus entry. (A) The N, S, E and M proteins are represented in their oligomeric state. N protein dimers bind +ssRNA, forming the nucleocapsid. The nucleocapsid is surrounded by the viral membrane that contains S, E and M proteins. The M protein is shown interacting with the S, E and N proteins. (B) Domain localization of the S protein, showing the S1 and S2 fragments; S1 contains the receptor binding domain (RBD), and S2 the fusion peptide (FP). (C) Angiotensin I converting enzyme 2 (ACE2) recognition by RBD, and the subsequent S proteolytic activation by a Furine-like protease or TMPRSS2. (D) Viral and host membrane fusion induction by the exposed FP.

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**Fig. 2.** SARS-CoV-2 genome organization. (A) Open reading frames (ORF) distribution in SARS-CoV-2 genome. (B) Non-structural proteins (nsp’s) distribution in orf1a and orf1ab, detailing multidomain organization of nsp3, nsp12, and nsp14; red and blue arrows indicate PL\textsubscript{pro} and M\textsubscript{pro} cleaving sites, respectively. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)
all +ssRNA viruses, the genome is copied to a reverse and complementary intermediate -ssRNA, which is copied back to a complete +ssRNA genome and several shorter sub-genomic +ssRNA, that are used as additional templates to translate high amounts of certain structural and accessory proteins, necessary during the viral assembly (Wu et al., 2020b). CoVs require the RdRp for replication, represented by the multi-protein complex nsp12-nsp8-nsp7. The nsp12 has the RdRp activity, nsp8 is the putative primase, while nsp7 acts as a co-factor (Peng et al., 2020). CoVs possess a helicase (nsp13) that unwinds structured or double-strand RNA (dsRNA) to allow the RdRp to proceed unhindered, thus achieving high efficiency and accuracy in RNA replication; they also have a 3'-5' exoribonuclease (ExoN, nsp14) that is capable of proofreading activity during RNA synthesis, thus lowering the rate of nucleoside misincorporation, while also enhancing resistance to nucleoside analogs (Bouvet et al., 2012; Jia et al., 2019) (Fig. 3).

Additionally, for the viral RNA to be translated efficiently and to avoid quick degradation, CoVs have poly-adenylation [poly(A)] and capping modifications, much like host mRNAs. The poly(A) modification is apparently catalyzed by nsp8, which has TAT activity (Tvarogova et al., 2019). The methyl capping consists of a sequence of reactions: 1) the terminal γ-phosphate is removed from the 5′-triphosphate end of RNA by a RNA 5′-triphosphatase (RTPase), probably nsp13; 2) an uncharacterized RNA guanylyltransferase (GTase) adds a GMP molecule to the 5′RNA, resulting in the formation of GpppN-RNA; 3) the GpppN-RNA is methylated at the N7 position of the guanosine by an N7-methyltransferase (N7-MTase, nsp10 dimer), yielding m7GpppN (cap-0); 4) finally, a 2′-O-MTase (nsp10 dimer) methylates the 2′-O position of the first nucleotide’s ribose of m7GpppN, yielding m7GpppNm (cap-1) (Snijder et al., 2016) (Fig. 4).

The last step in the viral cycle is the assembly of the viral particles and the release of the viral progeny. During this step, several N proteins bind and compact each copy of the genomic +ssRNA, forming the nucleocapsid. Proteins S, E, and M, accumulated in the ER-Golgi intermediate compartment (ERGIC) (Knoops et al., 2008), bind the nucleocapsid by direct M-N protein interactions (He et al., 2004), and then through the action of E and M proteins on the ERGIC membrane a viral particle is formed in a secretory compartment (Knoops et al., 2008). Finally, the viral progeny particle or virion is released by the exocytosis pathway (Fehr and Perlman, 2015), and the viral cycle is completed.

4. Druggability characterization of SARS-CoV-2 proteins

For each SARS-CoV-2 protein, we highlighted its relevant functional roles and main structural features, and then searched for druggable sites within the protein’s structure, analyzing the possible functional consequences of small-molecules binding to these sites. To find and assess the targetability of potential binding sites (including allosteric ones), we used FTMap to identify binding hot-spots on all available (or homology modeled) SARS-CoV-2 structures. FTMap samples a library of small organic probes of different molecular properties through docking, thus identifying consensus sites (CSs) or hot-spots (see Computational Methods for details). Hot-spots are small regions within a binding site considered to strongly contribute to the ligand-binding free energy. This analysis was complemented by the use of ICM Pocket Finder, and by the detection of cryptic sites, which are those usually “hidden” sites in unligated structures, but present only in ligand-bound structures. The strength of a CS is related to the number of fragment-like probe clusters it contains. A strong CS and nearby hot-spots define a potential binding site (Kozakov et al., 2015a), which may be considered druggable if conditions on the number of probe clusters within the primary CS and

![Fig. 3. Representative steps during the synthesis of a complementary RNA strain. Priming of the complementary strain, in this case, a -ssRNA, is catalyzed by nsp8 (red letters and arrows). Primer dependent RNA extension, catalyzed by nsp7-nsp8-nsp12 complex (blue lines and arrows). Magenta asterisk (*) and arrow represent a mis-incorporated nucleotide or nucleotide analog, and nonobligate RNA chain termination, respectively. Orange lines and arrows represent nucleotides forming dsRNA or structured RNA, and extension inhibition, respectively. Light green arrow represents nucleotide excision by nsp14 3′-5′ exonuclease (ExoN). Dark green arrow represents the unwinding activity of nsp13 helicase. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)](image-url)
the presence of other hot-spots in the vicinity are satisfied (see Computational Methods); these conditions mean that a drug-like molecule could bind the receptor with at least micromolar affinity (Kozakov et al., 2015a). It is clear that this assessment of druggability does not necessarily mean that a ligand binding at that site will actually exert an observable biological effect. We also summarized current drug repurposing efforts on the two cysteine proteases and the polymerase, but only including compounds with some kind of experimental validation.

4.1. Non-structural proteins

4.1.1. Non-structural protein 3 (nsp3)

Nsp3 is the largest SARS-CoV-2 protein, with 1945 amino acids. It has at least seven tandem macrodomains involved in different functions. These nsp3 domains are listed in Table S1, and each one is analyzed in the subsequent sections.

4.1.2. The papain-like protease (PLpro, nsp3 domain)

The inhibition of viral proteases might not only be important to block viral polyproteins, but it would be a way to interfere with viral-induced immunosuppression processes, since these proteases also act on host proteins, and are partly responsible for the host cell shutdown. The PLpro has a ubiquitin-like subdomain (Ub12) located within its N-terminal portion, which is not necessary for PLpro activity in SARS (Clasman et al., 2017), but it is probably involved in Nuclear Factor kappa-light-chain-enhancer of activated B cells (NFκB) signaling, affecting sensibility to interferon (IFN) (Friedman et al., 2009). Through recognition of the consensus cleavage LXGG motif, PLpro hydrolyzes the peptide bond on the carboxyl side of the Gly at the P1 position, thus releasing nsp1, nsp2, and nsp3 proteins. In the case of SARS-CoV, in vitro studies have also shown that this protease has two other proteolytic activities, hydrolyzing ubiquitin (Ub) and ubiquitin-like protein Interferon-Stimulated Gene product 15 (ISG15), from cellular proteins such as the Interferon Regulatory Factor 3 (IRF3), thus suppressing the host’s innate immune response (Matthews et al., 2014).

The crystal structures of SARS-CoV-2 PLpro in complex with inhibitors VIR250 and VIR251 covalently attached to the catalytic C111 were solved at 2.8 Å (VIR250, PDB 6WUU) and 1.65 Å (VIR251, PDB 6WX4), respectively (Figs. 5A and S1). These inhibitors displayed similar activities towards SARS-CoV and SARS-CoV-2 PLpro’s, but a weaker activity towards the MERS-CoV PLpro. Two wild type apo structures (PDBs 6W9C and 6WZU, at 2.7 Å and 1.8 Å, respectively), and the C111S apo structure (PDB 6WHR, at 1.6 Å) have overall main chain RMSD values of ~0.6 Å compared to the lowest resolution PLpro-VIR251 structure, displaying a partial collapse of the catalytic binding site. SARS-CoV-2 PLpro was crystallized also in complex with Ub at 2.7 Å (PDB 6XAA), and with the C-terminal part of ISG15 at 2.9 Å (PDB 6X9A) (Fig. S1). Both structures are very similar to 6WX4, with main chain RMSD values of ~0.5 Å (all available PLpro structures are listed in Table S1).

In agreement with another work (Freitas et al., 2020), it was shown that while SARS-CoV-2 PLpro retained de-ISGylating activity similar to its SARS-CoV counterpart, its hydrolyzing ability of diUbK48 was comparatively decreased (Rut et al., 2020). Interestingly, SARS-CoV-2 PLpro displays a preference for ISG15s from certain species including humans (Freitas et al., 2020). It has yet to be determined whether this functional difference has any implication in the ability of SARS-CoV-2 to evade the human innate immune system.

The catalytic site (Fig. 5) would clearly be the first choice for drug discovery (see below an account of successful cases targeting this site). Using FTPMap, a borderline druggable binding site delimited by residues R166, L185, L199, V202, E203, M206-M208, and K232, was identified; a small-molecule binding to this site might interfere with Ub binding, but not with ISG15 binding (Fig. 5B). A small hot-spot could also be found limited by amino acids P59, R65, and T74-F79. This site is close to the Ub S2 binding site, and from the structural model of the SARS-CoV-2 PLpro-diUbK48 (generated using the corresponding SARS-CoV structure (PDB 5E6J)), a small-molecule binding to it might interfere with Ub binding. Two other alternative potential binding sites were identified in PLpro, delimited by residues: s) S213-E215, K218, Y252, L254, K255, T258-T260, V304, Y306, and E308; ii) L121-I124, L126-F128, L133, Q134, Y137, Y138, and R141 (Fig. S1). These sites are far from the catalytic, Ub1-, and Ub2-binding sites, so that, even in the case a small-molecule could bind to them, further evidence would be needed to determine whether those amino acids have a functional role, and to assess whether those sites could display allosteric modulation potential.

Considering that the residues lining the catalytic sites of SARS-CoV and SARS-CoV-2 PLpro are identical, Freitas et al. evaluated naphthalene-based SARS-CoV non-covalent inhibitors on SARS-CoV-2, and found that two compounds, GRL-0617 and 6, exhibited IC50 values of 2.4 μM and 5.0 μM, respectively (their corresponding values in SARS-CoV PLpro were 600 nM and 2.6 μM), displaying EC50 values of 27.6 and 21.0 μM, respectively, in antiviral activity assessment, with no cytotoxicity in cell cultures (Freitas et al., 2020). The characterization of the non-covalent inhibition between these ligands and SARS-CoV-2 PLpro was performed using a homology model based on SARS-CoV PLpro bound to GRL-0617 (PDB 3E9S). It should be stressed that a similar approach using MERS-CoV inhibitors (Lee et al., 2019) would not be so straightforward, considering the lower conservation of the catalytic binding site at the sequence level. It was also shown that the synthetic organoselenium drug molecule ebselen, which displays anti-inflammatory, anti-oxidant and cytoprotective activity in mammalian cells, covalently inhibits the enzymatic activity of SARS-CoV-2 PLpro, with an IC50 ~ 2 μM, exhibiting a weaker activity
against its SARS-CoV counterpart (Welgarz Tomczak et al., 2020b). In a follow-up contribution, ebselen-derivatives were identified, displaying lower inhibition constants, in the range of 250 nM (Welgarz Tomczak et al., 2020b). Klemm et al. showed that benzodioxolane analogs against its SARS-CoV counterpart (Welgarz Tomczak et al., 2020b). In a follow-up contribution, ebselen-derivatives were identified, displaying lower inhibition constants, in the range of 250 nM (Welgarz Tomczak et al., 2020b). Klemm et al. showed that benzodioxolane analogs

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4.1.3. Main protease (Mₚᵣₒ, nsp5)

The SARS-CoV-2 Mₚᵣₒ is a cysteine protease involved in most cleavage events within the precursor polyproteins, beginning with the autolytic cleavage of itself from pp1a and pp1ab (Fig. 2). The vital functional role of Mₚᵣₒ in the viral life cycle makes it an attractive antiviral target.

The active form of Mₚᵣₒ is a homodimer containing two protomers composed of three domains each (Jin et al., 2020a): domain I (residues F8–Y101), domain II (residues K102–P184), and domain III (residues T201–V303); domains II and III are connected by a long loop region (residues F185–I200). The Mₚᵣₒ has a Cys-His catalytic dyad (C145–H41), and the substrate binding site is located in the cleft between domains I and II (Fig. S2). The superposition of 12 crystal structures of Mₚᵣₒ from different species (Jin et al., 2020a) showed that the helical domain III and surface loops display the largest conformational variations, while the substrate-binding pocket is highly conserved among CoVs.

Crystal structures of SARS-CoV-2 Mₚᵣₒ were determined in complex with the Michael acceptor covalent inhibitor N3 (Fig. 6A) at 2.1 Å resolution (PDB 6LU7), and 1.7 Å (PDB 7BQY), respectively (an up-to-date detailed list of Mₚᵣₒ solved structures is shown in Table S2). N3 inhibits the Mₚᵣₒ from multiple CoVs, and has been co-crystalized with Mₚᵣₒ in SARS-CoV (PDB 2AMQ), Infectious Bronchitis Virus (IBV) (PDB 2Q6F), Human coronavirus HKU1 (HCoV-HKU1) (PDB 3D23), Feline Infectious Peritonitis Virus (FIPV) (PDB 5EU8), Human coronavirus NL63 (HCoV-NL63) (PDB 5GWY), Porcine Epidemic Diarrhea Virus (PEDV) (PDB 5GZW), and Mouse Hepatitis Virus (MHV) (PDB 6JU). In the SARS-CoV-2 Mₚᵣₒ-N3 complexes, the Sγ atom of C145 forms a covalent bond with the Cj atom of the vinyl group of N3 (see Table S2 for available structures solved in the apo form, and with covalent and non-covalent inhibitors; more than a hundred crystal structures with bound fragment-like molecules are not included).

The primary choice for drug discovery would be the catalytic site. Zhang et al. crystallized SARS-CoV-2 Mₚᵣₒ in its apo form (PDB 6Y2E) to guide the design of a novel α-ketoamide inhibitor displaying an IC₅₀ = 0.67 μM; crystal structures of SARS-CoV-2 in complex with this inhibitor were determined in the monocular and orthorombic form (PDBs 6Y2F and 6Y2G, respectively). The structure of Mₚᵣₒ in complex with anti-neoplastic drug carmofur, which inhibits viral replication in cells with EC₅₀ = 24.3 μM, was determined at 1.6 Å resolution (Jin et al., 2020b); the carbonyl group of carmofur attaches covalently to catalytic C145 and the fatty acid tail occupies the S2 subsite. Anti-inflammatory ebselen has also shown a promising inhibitory effect in Mₚᵣₒ and reduction of the virus titer in cell culture (Jin et al., 2020a). Starting with the substrate-binding site of the SARS-CoV Mₚᵣₒ, Dai et al. designed and synthesized two novel covalent inhibitors of the SARS-CoV-2 Mₚᵣₒ, displaying anti-SARS-CoV-2-infection activity in cell culture with EC₅₀ values of 0.53 μM and 0.72 μM, respectively, with no significant cytotoxicity (Dai et al., 2020). Some known protease inhibitors have been identified in silico and tested in vitro on Mₚᵣₒ, such as the HIV protease inhibitors ritonavir, nelfinavir, saquinavir, and atazanavir (Fintelman-Rodrigues et al., 2020); the hepatitis C virus (HCV) protease inhibitor boceprevir, the broad-spectrum protease inhibitor GC-376, and calpain inhibitors II and XII were also shown to inhibit viral replication by targeting the Mₚᵣₒ (Ma et al., 2020).

In addition to the catalytic site, we identified a cryptic site with a CS within 5 Å defined by residues T199, Y237, Y239, L271, L272, G275, M276, and A285-L287; this site is not too distant from the partner protomer, and as such should be explored whether a molecule binding to it might preclude dimer formation (Fig. 6B). A second borderline druggable site delimited by residues Q107-Q110, V202, N203, H246, I249, and T292-F294 was also identified. This site lies on the opposite...
4.1.4. The RNA polymerase complex (nsp12-nsp7-nsp8)

In +ssRNA viruses, the synthesis of RNA is catalyzed by the RdRp, in a primer-dependent manner. In SARS-CoV-2 and other CoVs, the RdRp complex consists of a catalytic subunit (nsp12), and co-factors nsp7 and nsp8 (Gao et al., 2020; Kirchdoerfer and Ward, 2019).

The SARS-CoV-2 RdRp experimental structures contain two nsp8 molecules: one forming a heterodimer with nsp7 (nsp8-2), and a second one bound at a different site (nsp8-1), in a similar fashion as in SARS-CoV (Kirchdoerfer and Ward, 2019) (cf. Table S3 for a list of the available RdRp complex experimental structures). Interaction with nsp7 and nsp8 provides stability to nsp12 (Peng et al., 2020), consistent with the observation that nsp12 in isolation displays little activity, while the presence of nsp7 and nsp8 enhances template binding and processivity (Kirchdoerfer and Ward, 2019; Yin et al., 2020). The overall structure of the SARS-CoV-2 RdRp complex is very similar to that of SARS-CoV, with a Cα RMSD of ~0.8 Å, consistent with the high degree of sequence similarity (nsp7, 99%; nsp8, 97%; nsp12, 97%). Although the amino acid substitutions are not located in the catalytic site, the SARS-CoV-2 RdRp complex displays a 35% lower efficiency for RNA synthesis than its SARS-CoV counterpart; this lower efficiency is due to changes restricted to nsp8 and nsp12 only (Peng et al., 2020).

The nsp12 contains a right-hand RdRp domain (residues L366-F932) – a conserved architecture in viral RdRps and an N-terminal nido-virus RdRp-associated nucleotidyl-transferase (NiRAN) domain (residues D51-R249); these two domains are linked by an interface domain (residues H816-E932) (Fig. 7A). An N-terminal β-hairpin subdomain (residues A250 to R365) interacts with nsp7 through its phosphate-ribose backbone, especially involving the residues D29-K50 establishes close contacts with the NiRAN domain and the palm subdomain, and contributes to stabilize the structure. As in the corresponding SARS-CoV nsp12 (Kirchdoerfer and Ward, 2019), the integrity of the overall structure is maintained by two zinc ions that are present within metal-binding sites composed by residues H295–C301–C306–C310 and C487–H642–C645–C646, respectively.

The RdRp-RNA structures show that RNA mainly interacts with nsp12 through its phosphate-ribose backbone, especially involving the Z-OH groups. Except for three specific and localized structural changes (Yin et al., 2020), the overall structures of the apo and RNA-bound complexes are very similar, displaying a main chain RMSD value of ~0.5 Å. The absence of relevant conformational rearrangements from the apo state implies that the enzyme can start to function as replicase upon RNA binding, which also correlates with its high processivity. The RNA binding residues are highly conserved across viral polymerases (teVelthuis, 2014). It should be noted that in RdRp structures with longer RNA products (PDBs 6YYT, 26 bases, and 6XEZ, 34 bases), the long α-helical extensions of both nsp8 molecules form positively charged ‘sliding poles’ along the exiting RNA, in what could be a supporting scaffold for processing the large CoV genome. Since these nsp8 extensions are mobile in free RdRp (Gao et al., 2020; Kirchdoerfer and Ward, 2019), it is possible that they adopt an ordered conformation upon dsRNA exit from nsp12.

Remdesivir, a drug designed for Ebola virus, has received a great deal of attention as a RdRp inhibitor and potential treatment for COVID-19 (Wang et al., 2020a). Remdesivir is a prodrug that is converted into its active triphosphate form [remdesivir triphosphate (RTP)] by host enzymes. Two cryo-electron microscopy structures of the SARS-CoV-2 RdRp complex have been solved, one in the apo form, and the other in complex with a template-primer RNA and with remdesivir covalently linked (teVelthuis, 2014). It should be noted that in RdRp structures with longer RNA products (PDBs 6YYT, 26 bases, and 6XEZ, 34 bases), the long α-helical extensions of both nsp8 molecules form positively charged ‘sliding poles’ along the exiting RNA, in what could be a supporting scaffold for processing the large CoV genome. Since these nsp8 extensions are mobile in free RdRp (Gao et al., 2020; Kirchdoerfer and Ward, 2019), it is possible that they adopt an ordered conformation upon dsRNA exit from nsp12.

Fig. 7. The SARS-CoV-2 RdRp complex (nsp12-nsp8-nsp7). (A) Structure of the nsp12-nsp7-nsp8 complex. The channel in the middle corresponds to the catalytic active site. Color code: nsp7, white ribbon; nsp8-2, light blue ribbon; nsp8-1, yellow ribbon. The nsp12 domains are colored as follows: palm, yellow; fingers, tan; palm, red; interface, pale green ribbon; NiRAN, magenta ribbon. (B) RdRp complex bound to RNA. Nsp12 is displayed as a green surface. Color code: Primer RNA, blue; template RNA, red; nsp7, yellow ribbon; nsp8-1, grey ribbon. (C) Molecule of ADP-Mg2+ within the NiRAN domain of nsp12. Interacting residues are shown, in what may constitute a drug-binding site. (D) Target site in nsp8 (light yellow surface). The predicted binding site is represented using a blue mesh representation, and nsp12 is shown as grey ribbon. Nsp12 residues N386–K391 are displayed (though not labeled) to highlight that a small molecule binding to these potential sites might interfere with PPIs. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)
form of remdesivir (RMP) is added (at position i) into the primer-strand (Yin et al., 2020), thus causing termination of RNA synthesis at position i+3, in SARS-CoV-2, SARS-CoV, and MERS-CoV (Gordon et al., 2020a). Favipiravir (also called avifavirus), used in Hepatitis C Virus (HCV) treatment, is also a promising drug (Wang et al., 2020a). Other antivirals, such as sofosbuvir, alavudine, tenofovir alafenamide, AZT, abacavir, lamivudine, emtricitabine, carbovir, ganciclovir, stavudine, and entecavir, are also incorporated by SARS-CoV-2 RdRp and block replication in vitro (Chien et al., 2020). Another type of broad spectrum ribonucleoside analog, β-D-N4-hydroxyctydine (EIDD-1931), has been shown to inhibit SARS-CoV-2, SARS-CoV and MERS-CoV in cell culture by increasing the mutation transition rate, probably exceeding the proofreading ability granted by the ExoN (Sheahan et al., 2020).

In the RdRp-nsp13 complex (PDB 6XEZ), the NiRAN domain is occupied by an ADP-Mg\textsuperscript{2+} molecule (Fig. 7C). While the target of the NiRAN nucleotidyltransferase activity is unknown, the activity itself is essential for viral propagation (Lehmann et al., 2015). Thus, the ADP-binding site might constitute an interesting druggable site in nsp12, and it should be further assessed whether a small-molecule binding to it would interfere with the viral cycle.

The design of non-nucleoside inhibitors calls for the analysis of alternative disruptive sites within the RdRp complex, considering that the assembly of the nsp12-nsp7-nsp8 complex is needed for RNA synthesis. We used FTMap to identify druggable sites on PPI interfaces in nsp12, nsp7, and nsp8. Several hot-spots were identified on nsp8, three of which formed a potential druggable site, outlined by residues P121, A125, K127–P133, T137, T141, F147, and W154 (Fig. 7D). This site lies within the nsp12-nsp8 interface, and a molecule binding to this site could interfere with the interaction of the β-strand N386–K391 of nsp12 with nsp8 (Fig. 7D). A borderline druggable target site was identified on nsp12, lined up by residues L270, L271, P323, T324, F326, P328–V330, R349, F396, V398, M666, and V675 (Fig. S3), where a molecule binding to it might interfere with nsp8 binding by clashing with its segment V115–II19. On nsp7, a borderline druggable site was identified within the PPI interface with nsp8-2, defined by residues K2, D5, V6, T9, S10, V12, F49, and M52-S55.

In addition to its functionality within the RdRp complex, the nsp7-nsp8 hexadecamer has de novo initiation of RNA synthesis capability, also known as primase activity, while nsp8 has also displayed TATase activity. Analysis of the SARS-CoV nsp7-nsp8 hexadecameric structure (PDB 2AHM) shows several sites which could be targeted to interfere with its primase activity, making impossible for nsp12 to extend the complementary strand due to the lack of primer. Moreover, it is also possible that small-molecules could interfere with the conformational dynamics needed to interact within the RdRp complex. However, further structural and computational studies (such as molecular dynamics, MD) would be needed to confirm these hypotheses.

4.1.5. Helicase (nsp13)

The SARS-CoV-2 nsp13 possesses helicase activity, thus playing a key role in catalyzing the unwinding in the 5′ to 3′ direction of dsRNA or structured RNA into single strands. It has been demonstrated that in SARS-CoV this happens in an NTP-dependent manner (Jia et al., 2019). Additionally, it has been shown that nsp13 has RNA 5′-triphosphata (RTPase) activity (which may be the first step for the formation of the 5′ cap structure of viral RNAs). The helicase-associated RTPase and RTFase activities share a common active site, both in SARS-CoV (Ivanov et al., 2004) and the human CoV 229E (HCoV-229E) (Ivanov and Ziebuhr, 2004).

The crystal structure of the SARS-CoV-2 nsp13 has been solved at 1.9 Å (PDB 6ZSL). A structure solved by cryo-EM of nsp13 in complex with RdRp (nsp12-nsp8-nsp7) is also available (PDB 6XEZ). Nsp13 has the form of a triangular pyramid with five domains. At the top, the N-terminal zinc binding domain with three zinc fingers (A1-S100) is connected with the stalk domain (D101-G150); then, at the base of the pyramid, three domains (1B, I151-E261; 1A, F262-R442; 2A, R443-N596) form the triangular base (Fig. S4). Compared to its SARS-CoV and MERS-CoV counterparts, the SARS-CoV-2 nsp13 shares a 99.8% (100%), and 71% (82%) sequence identity (similarity), respectively. The nsp13 structures are also very similar, with backbone RMSD values of ~1.9 Å between SARS-CoV-2 nsp13 and the corresponding SARS-CoV nsp13 (PDB 6UYT) and MERS-CoV (PDB SWWP).

In the SARS-CoV nsp13, six residues were identified as being involved in NTP hydrolysis (K288, S289, D374, E375, Q404, and R567), and mutations of any of these residues to alanine resulted in high unwinding deficiency and decreased ATPase activity (Jia et al., 2019). Moreover, it was also shown in the same study that, for all six mutants, changes in helicase activity are consistent with changes in their ATPase activity, thus demonstrating that nsp13 performs its helicase activity in an NTP-dependent manner. As stated above, this site would also correspond to the RTPase activity. The SARS-CoV-2 nsp13 structure PDB 6XEZ features an ADP-Mg\textsuperscript{2+} molecule in the vicinity of those residues (Fig. 8A), suggesting that targeting this site may interfere with both NTPase or RTPase functions.

The zinc binding and stalk domains are important for the helicase activity in SARS-CoV (Jia et al., 2019). Mutations to alanine of the zinc binding domain residue N102 (which interacts with the stalk domain) and of the stalk residue K131 (which interacts with the 1A domain) resulted in decreased helicase activity. This appears to be a top-to-bottom signaling system, and it is difficult to figure out a way to interfere in this process with small molecules, with the information available up to today. The zinc binding domain might be targeted with metal chelators such as bismuth complexes.

Based on an interaction model of dsRNA with SARS-CoV nsp13, it was hypothesized that residues 176–186 (1B domain), 209–214 (1B domain), 330–350 (1A domain) and 516–541 (2A domain) constitute a probable nucleic acid binding region (Jia et al., 2019). While there is no CoV nsp13 structure with a nucleic acid substrate bound, we performed a crude model of SARS-CoV-2 nsp13 in complex with RNA based on a Yeast-Upf1-RNA structure (PDB 2XZL), in a similar fashion as in an earlier work (Jia et al., 2019), and in agreement with a recent model (Chen et al., 2020) (Fig. 8B). Using FTMap, two potential borderline druggable sites were identified delimited by amino acids P406–409, L412, T413, G415, L417, F422, N557, and R560, and K139, E142, C309, M378–D383, P408, and T410, which could interfere with RNA binding, according to our artificial model (Fig. 8B).

4.1.6. Exoribonuclease/guanine-N7 methyltransferase (nsp14-nsp10 complex)

While the RdRp complex allows the synthesis of large viral RNA due to an increased processivity conferred by the heteromeric complex, to achieve high accuracy, CoVs possess 3′-5′ exoribonuclease (ExoN) activity located in the N-terminal domain of nsp14, to proofread the viral genome synthesis. The ExoN excises mutagenic nucleotides mis-incorporated by the RdRp, thus conferring potential drug resistance to nucleoside analog inhibitors. It has been reported that the ExoN activity protects SARS-CoV from the effect of base analog 5-fluorouracil (Smith et al., 2013), and that guanosine analog ribavirin (Rbv) 5′-monophosphate is incorporated at the 3′-end of RNA by the SARS-CoV RdRp, but excised from RNA by the nsp14-nsp10 ExoN (to a lesser degree by the nsp14-ExoN alone), which could account for the poor effect of Rbv in treating CoV-infected patients (Al-Tawfiq and Memish, 2017). Since it was shown that SARS-CoV displays a significantly lower nucleotide incorporation fidelity in vitro than that of the Dengue RdRp (Ferron et al., 2018), it could be concluded that the low mutation rate of SARS-CoV is related to the ExoN activity. This shows that targeting the ExoN function could be an excellent strategy for the development of pan-CoV therapeutics, complementing existing RdRp inhibitors, like Remdesivir.

In fact, MVH lacking the ExoN activity was shown to be more susceptible to Remdesivir (Agostini et al., 2018).

The C-terminal domain of nsp14 functions as a guanine-N7 methyltransferase (N7-MTase), where S-adenosyl-L-methionine (SAM) is
demethylated to produce S-adenosyl-L-homocysteine (SAH), transferring the methyl group to the N7 position of the guanine in 5-demethylated RNA, and forming 7GpppN (cap-0). This capping is followed by a second methylation by the 2′-O-MTase (nsp16), forming 7GpppNm (cap-1) (Fig. 4). The capping structure is a protective and protosungal modification of the viral RNA, and blocking MTase activity may increase viral RNA decay and suppress viral RNA translation.

While there is no experimental structure available of SARS-Cov-2 nsp14, crystal structures of the SARS-CoV nsp10-nsp14 dimer are available, with nsp14 in complex with SAM at 3.2 Å (PDB 5C8T), with SAH and guanosine-P3-adenosine-5′,5′-triphosphate (GpppA) at 3.3 Å (PDB 5CBS), and in its unbound form (PDBs 5C8U and 5NFY, both at 3.4 Å). The sequence identity between SARS-Cov-2 and SARS-CoV nsp14 is 95% (98% similarity), with no gaps in the alignment and full conservation in all functionally relevant sites. SARS-CoV nsp10 and its SARS-CoV-2 counterpart share 98% sequence identity (similarity 100%). We thus built a homology model of the SARS-Cov-2 nsp10-nsp14 dimer using the corresponding SARS structure 5CBS as a template (see Methods); the missing S454-D464 segment within the template was included in the model, and optimized. Since there are no gaps, the numbering scheme of template and model coincides.

In nsp14, amino acids A1-C285 fold into the ExoN domain, and the N7-MTase function lies within amino acids D301-Q527; both domains are connected by a loop (amino acids F286-G300), and its abolition was shown to suppress the N7-MTase function in SARS-CoV (Chen et al., 2013) (Fig. 9A). The architecture of the catalytic core and active sites of the ExoN domain are similar to those of the DEDD superfamily exonucleases, though exhibiting some differences (Ferron et al., 2018; Ma et al., 2015). The catalytic residues D90, E92, E191, D273 and H268 ion at its active center. Mutating any of these residues to alanine impairs the ExoN activity or severely reduced the ability to degrade RNA in SARS-CoV (Ma et al., 2015). The ExoN domain also contains two zinc fingers, the first one comprising residues C207, C210, C226 and H229, and the second one in proximity to the catalytic site, comprising residues H257, C261, H264 and C279. In SARS-CoV, none of the mutants of zinc finger 1 could be expressed as soluble proteins, thus revealing its importance in protein stability, and mutants of zinc finger 2 containing two zinc fingers, the first one comprising residues C207, C210, C226 and H229, and the second one in proximity to the catalytic site, comprising residues H257, C261, H264 and C279. In SARS-CoV, none of the mutants of zinc finger 1 could be expressed as soluble proteins, thus revealing its importance in protein stability, and mutants of zinc finger 2 could not be expressed in SARS-CoV-2.

As it has been pointed out in SARS-CoV (Ferron et al., 2018; Ma et al., 2015), nsp14 has an atypical MTase fold, with an additional α-helix in the last 12 amino acids which stabilizes the neighboring environment; deletion of this helix has been shown to decrease or abolish the MTase activity of nsp14. A third zinc finger is also present in this domain formed by C452, C477, C484 and H487, but it is distant from the active site, and mutations of amino acids from the zinc finger 3 have a very weak effect on nsp14 activity.

Fig. 8. The SARS-CoV-2 helicase/NTPase/RTPase (nsp13). (A) The ADP-Mg\(^{2+}\) bound within nsp13. This site has been identified as being involved in NTP hydrolysis in SARS-CoV, and could constitute a druggable site. (B) Two potential border-line druggable binding sites identified in nsp13. The structure of RNA (blue) was modeled based on the yeast Upf1-RNA complex structure (PDB 2XZL). Nsp13 is represented as a green ribbon, but nsp13 domains 1A and 2A are displayed as grey and tan molecular surfaces, respectively. The potential sites are shown in yellow. Based on our model, molecules binding to these sites might interfere with RNA binding. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)
marginal effect on MTase activity (Ma et al., 2015). But it has been hypothesized that it might be important in binding with the nsp16-nsp10 dimer to accomplish the second methylation for completion of the capped structure.

The SARS-CoV-2 N7-MTase domain in complex with methyl donor SAM is shown in Fig. 9B; the binding site residues are fully conserved, thus the contacts are similar to the SARS-CoV nsp14-SAM complex. Methyl acceptor GpppA binds close to methyl donor SAM (cf. PDB 5C8S) to facilitate methyl transfer. Comparing the N7-MTase catalytic sites of SARS-CoV nsp14 structures bound to SAM (5C8T), and to SAH-GpppA (5C8S), no significant structural changes are found, thus supporting the hypothesis that ligand binding sites are pre-formed.

The hinge region (amino acids F286-G300, Fig. 9A) is conserved across CoVs, thus suggesting it might have a functional role. In SARS-CoV, lateral and rotational movements of the C-terminal domain relative to the N-terminal domain of up to 13 Å have been observed (Ferron et al., 2018), and crystallographic and SAXS data have shown that nsp14 undergoes important conformational changes, where the hinge might act as a molecular switch; moreover, hinge residues Y296 and P297 are essential for ExoN activity in SARS-CoV.

In fact, nsp14 is involved in two processes that use RNA substrates in different ways: a newly synthesized RNA strand with a mismatch should be translocated by the polymerase complex to the ExoN catalytic site of nsp14, whereas during replication, the 5′-mRNA should go into the catalytic tunnel of the N7-MTase for methyl capping. These results could be rationalized in terms of nsp14 flexibility. Moreover, the mutual dependence of the ExoN domain on N7-MTase function (Chen et al., 2013), and of the N7-MTase domain on ExoN activity (Ferron et al., 2018) have been established. Ferron et al. further showed that both the ExoN domain (excluding the first 71 residues) and the N7-MTase domain interact with nsp12-RdRp (Ferron et al., 2018), a biologically relevant interaction due to the possibility of tandem associated activities that makes RNA replication more efficient.

Considering the structural and functional features of the SARS-CoV-2 nsp10-nsp14 dimer just discussed, the following small-molecule targeting strategies could be suggested:

- The SAM binding site is an attractive target to develop CoVs inhibitors using small-molecules that could preclude SAM or GpppA binding, thus suppressing the N7-MTase activity of nsp14. In fact, the N7-MTase catalytic tunnel lies close to the SAM binding site, and blocking it would also preclude mRNA binding.
- A potential druggable (allosteric) site was identified using FTMap and ICM Pocket Finder defined by residues R81-A85, L177-D179, Y296–I299, N408, L409, L411, and V421. This site lies in the vicinity of hinge residues Y296 and P297 (Fig. 9C). A small-molecule binding within this site could interfere with the dynamic behavior of nsp14 and its associated conformational changes.
- There is a small hot-spot in nsp14 (lined up by residues F60, M62, L192, M195, and K196) in the vicinity of a cysteine site, in which a molecule binding in this region would clash with nsp10 binding.
- Considering the high sequence identity between the SARS-CoV and SARS-CoV-2 nsp10-nsp14 dimers, and that the contact residues are fully conserved, blocking the nsp10-nsp14 interaction to decrease or abolish full ExoN activity could be a valid strategy against CoV diseases. A potential borderline druggable site on the PPI surface of nsp10 with nsp14 and lined up by residues T5, E6, N40, A71, S72, C77, L80, I92, K95, and Y96 was identified using FTMap; this site is defined by two hot-spots separated by ~9 Å, which could impose a limit on the expected affinity of a potential ligand (Kozakov et al., 2015b). However, it should also be mentioned that in the absence of nsp10, the nsp14 ExoN catalytic site partially collapses and its activity decreases (Bouvet et al., 2012); thus, it would be interesting to further explore the behavior of this site using MD simulations.
- A druggable site was identified within a region of a high cryptic site score in nsp10 (Fig. 9D). This site does not overlap with the nsp14-nsp16 PPI interface, and while its functional role is uncertain, it may have allosteric modulation potential.

4.1.7. RNA nucleotide-2′-O-methyltransferase complex (nsp16-nsp10)

The nsp 16 possesses a SAM-dependent RNA 2′-O-MTase activity that is capable of cap-1 formation. It adds a methyl group to the m7GpppN previously capped by nsp14 (cap-0) to form m7GpppNm (cap-1), by methylating the ribose of the first nucleotide at position 2′-O (Fig. 4). Like nsp14, nsp16 uses SAM as methyl donor (Decroly et al., 2008). In SARS-CoV, nsp16 requires nsp10 to execute its activity, since nsp10 is necessary for nsp16 to bind both m7GpppA-RNA and SAM (Chen et al., 2011); moreover, nsp16 was found to be an unstable protein in isolation, and most of the disruptions in the interface of nsp16-nsp10 eliminate the methyltransfer activity (Romas-Lemus et al., 2020). In humans, most mRNAs include the cap-1 modification; while cap-0 appears to be sufficient to recruit the entire translational machinery, cap-1 modification is necessary to evade recognition by host RNA sensors, such as the Retinoic acid-Inducible Gene I (RIG-I), MDA-5, and IFN-induced proteins with tetratricopeptide repeats (IFIT), and to resist the IFN-mediated antiviral response (Devarkar et al., 2016).

Several crystal structures of the nsp16-nsp10 heterodimer have been recently solved (cf. Table S4) (Fig. 10A). SARS-CoV-2 nsp10 shares 99% and 59% sequence identity with SARS-CoV and MERS-CoV, respectively. Similarly, nsp16 is highly similar to SARS-CoV (95% sequence identity), but only 66% identical to MERS-CoV. Both proteins interact through a large network of hydrogen-bonds, water-mediated interactions, and hydrophobic contacts (Kraficikova et al., 2020). The high conservation of nsp10 and nsp16 sequences, and the complete conservation of catalytic and substrate-binding residues strongly support the idea that nsp16-mediated 2′-O-MTase mechanism and functionality are highly conserved in CoVs. Nsp10 exhibits two zinc fingers: the first one is coordinated by C74, C77, H83 and C90, and the second one is coordinated by C117, C120, C128, and C130. The zinc fingers residues are 100% conserved across β-CoVs, highlighting the relevance of this motif in the replication process.

Most nsp16-nsp10 structures feature SAM (or the close analog pan-MTase inhibitor sinefungin) in their substrate binding site, being coordinated by residues N43, Y47, G71, A72, S74, G81, D99, N101, L100, D114, and M131, together with few water molecules (Fig. 10B). All these residues are also 100% conserved in SARS-CoV. Some structures also include mGpppA within the catalytic site (PDBs 6WQ3 and 6WRZ in presence of SAH, and 6WKS and 6WVN in presence of SAM). The nucleotide binding site is surrounded by flexible loops comprised of amino acids D26-K38, M131-N138, and the highly conserved KDKE motif (K46, D130, K170, E203) for methyl-transfer, which is found in many 2′-O-MTases (including the phylogenetically more distant MERS-CoV), is also present (Fig. 10B). The MTase active site is clearly an attractive target for antiviral drug discovery, but the structural conservation among CoVs and host cellular MTases might pose a challenge for the development of a specific compound.

Using FTMap, a small hot-spot was found in the vicinity of residues L57, T58, A188, C209, N210, and S276, within an extended cryptic site on the surface of nsp16 (Fig. 10D). Interestingly, recent crystal structures of the nsp16-nsp10 dimer feature small-molecules bound within that site, such as adenosine (PDB 6WKS), 2′-O-(methyl)-adenosine (PDB 6Y8Z), 2′-O-(methyl)-β-D-fructopyranose (PDB 6W4H), 7-methylguanosine-5′-triphosphate (PDBs 6WVN and 6WRZ), and 7-methylguanosine-5′-diphosphate (PDB 6WQ3) (Fig. 10C). This site lies on the opposite side of the catalytic site, 25 Å away from it, in what could thus be an allosteric site (Viswanathan et al., 2020). Further studies would be necessary to clarify the function of this binding site, and its impact and feasibility in terms of druggability.

Considering that m7GpppA acts as an effective substrate of SARS-CoV nsp16-nsp10 (Züst et al., 2011), Chen et al. performed docking of
m7GpppGAAAAA and m7GpppAAAAAA within the SARS-CoV nsp16-nsp10 dimer (Chen et al., 2011), and found that while the first three nucleotides contact nsp16, the following ones were in contact with nsp10, which extends the positively charged area of the RNA-binding groove. Interestingly, all the residues defining this groove extension are also conserved in SARS-CoV-2. The extension to the RNA-binding site provided by nsp10 may serve to accommodate the RNA chain and stabilize the interaction between m7GpppA-RNA within the catalytic site, which could also be observed by superposing the nsp16 SAM site, which could also be observed by superposing the nsp16 SAM site. (B) Catalytic site of nsp16 with methyl donor SAM (light yellow carbons) and methyl acceptor m7GpppA (cyan carbons). The nucleotide binding site flexible loops (D26-K38, M131-N138) are colored in blue. The highly conserved KDKE motif (K46, D130, K170, E203) for methyl-transfer, found in many Z-O-MTases, is highlighted in magenta, and oxygen water molecules are displayed in red. (C) Binding hot-spot (transparent yellow) identified with FTMap in the vicinity of residues L57, T58, A188, C209, N210, and S276, on the surface of nsp16, within an extended cryptic site identified in the same region using Cryptosite (brown colored surface). Small-molecules bound within that site (taken from crystal structures) are also displayed (light yellow carbon atoms): [adenosine, 2-(n-morpholino)-ethanesulfonic acid, β-D-fructopyranose, 7-methyl-guanosine-5′-triphosphate, and 7-methyl-guanosine-5′-diphosphate]. This site lies on the opposite side of the catalytic site. ~25 Å away from it, in what could thus be an allosteric site. (D) Extension of the RNA groove in nsp16 towards nsp10. Five RNA nucleotides are shown (light yellow carbon atoms), which correspond to those of the human mRNA 2′-O-MTase (PDB 4N48), after structural superposition of the binding site residues. SAM is displayed in grey carbons. The consensus site identified with FTMap is shown in yellow mesh. Nsp16 and nsp10 are colored according to their electrostatic potential (blue, positively charged; red, negatively charged; white, neutral). (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

m7GpppGAAAAA and m7GpppAAAAAA within the SARS-CoV nsp16-nsp10 dimer (Chen et al., 2011), and found that while the first three nucleotides contact nsp16, the following ones were in contact with nsp10, which extends the positively charged area of the RNA-binding groove. Interestingly, all the residues defining this groove extension are also conserved in SARS-CoV-2. The extension to the RNA-binding site provided by nsp10 may serve to accommodate the RNA chain and stabilize the interaction between m7GpppA-RNA within the catalytic site, which could also be observed by superposing the nsp16 SAM binding site with that of the human mRNA 2′-O-MTase (PDB 4N48) (Krafickova et al., 2020) (Fig. 10D). Using FTMap, we identified a small borderline druggable site within nsp10, in the area where RNA would extend (Fig. 10D). This pocket lies within the RNA binding groove extension described above, and is lined up by the side chains of amino acids P37, T39, C41, K43, F68, A104, and P107 of nsp10. A molecule binding to it might interfere with extended RNA binding. It should be stressed that, due to its zinc fingers, nsp10 has the ability to bind poly-nucleotides non-specifically (Matthes et al., 2006).

It was also shown that 12-mer and 29-mer peptides extracted from the dimerization surface of SARS-CoV nsp10 (F68–H80 and F68–Y96, respectively) were found to inhibit the activity of nsp16 at IC50 ~ 160 μM (Ke et al., 2012); this suggests that using short peptides might be a possible strategy, considering there is 100% sequence conservation within the nsp10/nsp16 interface for SARS-CoV and SARS-CoV-2. Interfering with the nsp16-nsp10 dimer formation using small molecules might still be challenging, however, due to the large area of contact and the absence of buried pockets at the nsp10-nsp16 interaction surfaces. In spite of this, a cryptic site was identified on nsp16 in the region of the PPI interface with nsp10 (Fig. S5), close to where the 12-mer and 29-mer peptides would bind. However, no CSs were found near this cryptic site, so further studies are needed to confirm this site as a potential target to inhibit oligomerization.

**4.1.8. RNA uridylate-specific endoribonuclease (nsp15)**

The nsp15 harbors a nidoviral RNA uridylate-specific endoribonuclease (NendoU) that belongs to the EndoU family, whose activity is to cleave downstream of uridylate, releasing 2′-O-MTase (NendoU) that belongs to the EndoU family, whose activity is to cleave downstream of uridylate, releasing polyuridines produced during the priming of the poly(A) ssRNA in replication (Fig. 3), which helps to dampen dsRNA melanoma differentiation-associated protein 5 (MDAS)-dependent antiviral IFN responses (Hackbart et al., 2020). Clearly, small molecules blocking RNA catalytic sites, or interfering with nsp15 oligomerization, would inhibit the enzyme activity and would help trigger cellular antiviral mechanisms. The SARS-CoV-2 nsp15 structure displays an N-term oligomerization domain, a middle domain, and a C-term NendoU catalytic domain (Kim et al., 2020) (Fig. S6A). Nsp15 has been crystallized in its apo form (PDB 6VWW at 2.2 Å resolution), and in complex with citric acid (PDBs 6XDH at 2.35 Å, and 6W01 at 1.9 Å), uridine-3′-monophosphate (PDB 6X4I at 1.85 Å), uridine-5′-monophosphate (PDB 6WLC at 1.85 Å), drug Tipiracil (PDB 6WXC at 1.85 Å), and product di-nucleotide GpU (PDB 6X1B at 1.19 Å). All structures are very similar, with main chain RMSD values of less than 0.5 Å between any pair of them. Taking structure 6W01 as a reference, SARS-CoV-2 nsp15 exhibits ~0.5 Å RMSD with respect to its SARS-CoV counterpart (88% sequence identity, 95% similarity), and 1.1 Å with respect to MERS-CoV nsp15 (51% sequence identity, 65% similarity) (cf. Ref (Kim et al., 2020), for a detailed comparison of SARS-CoV-2 nsp15 with SARS-CoV and MERS-CoV structures).
The catalytic active site within the NendoU domain is formed by residues H235, H250, K290, V292, S294, T341, and Y343 (Fig. S6B) (also conserved in SARS-CoV and MERS-CoV), with H235, H250, and K290 as the proposed catalytic triad (Kim et al., 2020). The catalytic residues also form a druggable binding site, as can be confirmed by its shape and the crystal structure of drug Tipiracil in complex with nsp15 (PDB 6WXC). The catalytic activity has been observed to be metal-dependent in most NendoU proteins, although there could be exceptions in other nidoviruses (Nedialkova et al., 2009). Although no crystal structure has been found with a metal ion (nor complexed with RNA), a metal binding site required for maintaining the conformation of the active site and substrate during catalysis has been proposed, coordinated by the carboxylate group of D283, the hydroxyl group of S262, and the carbonyl oxygen of P263.

SARS-CoV-2 nsp15 folds into a hexamer (a dimer of trimers) (Kim et al., 2020), in agreement with an earlier work showing that SARS-CoV nsp15 conformationally exists as a hexamer (PDB 2RHB), and also suggesting oligomerization-dependent endonuclease activity (Guarino et al., 2005). The channel of the hexamer is ~10 Å wide, open from top to bottom, and the hexamer is stabilized by extensive contacts between monomers. As such, it might be disrupted or destabilized by mutations or small molecules (Kim et al., 2020). Using FTMap, we identified two potential druggable sites; the first one is delimited by residues K71, N74, N75, M272, S274–N278, I328, L346, and Q347 (Fig. S6C), and close to the binding area of another protomer within the hexamer (Fig. S6C); a small-molecule binding to this site might disrupt PPI; the second one, delimited by residues E69, K71, K90, T196, S198, L252, D273, K277, Y279, V295, and D297, is deep and rather buried, within an area with a higher-than-average cryptic site score and an opening towards the hexamer channel.

4.1.9. Non-structural protein 9 (nsp9)

Nsp9 acts as a hub that interacts with several viral components; it binds ssRNA, nsp8, and the N protein (Gordon et al., 2020b; Sutton et al., 2004). Proteomic experiments showed that SARS-CoV-2 nsp9 also binds to several nuclear pore proteins (Gordon et al., 2020b), a process well characterized for other viruses that affect nuclear shutting as part of their replicative and host shutdown mechanisms. SARS-CoV nsp9 exists as a homodimer in solution formed through the interaction between the GXXXG motifs in parallel α-helices of the opposite protomers (Sutton et al., 2004); mutations in any of the glycines inducing dimer disruption have been associated with impeded viral replication and reduced RNA binding in SARS-CoV (Miknis et al., 2009), and in porcine delta coronavirus (PDCoV) (Zeng et al., 2018). Thus, as it has been already suggested, disruption of the homodimeric interface could be an appealing strategy against CoV-associated diseases (Hu et al., 2017). Homologs of nsp9 have been found in other β-coronaviruses such as SARS-CoV, MERS-CoV, HCoV-229E, and the avian IBV, with different degrees of sequence similarity compared to SARS-CoV-2. The mechanism of RNA binding to nsp9 and how it enhances viral replication, are not yet fully understood, but apparently depend on several positively charged amino acids on the surface (Zeng et al., 2018). While it is not known whether SARS-CoV-2 nsp9 plays an analog role like its SARS-CoV counterpart, their high sequence identity (97%), and their strong structural similarity (see below), might suggest a highly conserved functional role.

The structure of the SARS-CoV-2 nsp9 homodimer has been solved at 2.0 Å (PDB 6WXD) and at 3.0 Å (PDB 6W4B); both structures share a backbone RMSD of 0.9 Å and 0.5 Å, respectively. The monomer has a backbone RMSD of 0.94 Å compared to that of SARS-CoV nsp9 (PDB 1QZB); the RMSD value decreases to 0.44 Å when considering α-helices and β-sheets only. The protomers interact mainly through van der Waals contacts of the backbone from the conserved GXXXG motifs within the opposed α-helices, near the C-terminus of the protein (Fig. S7A), as in other CoVs (Figloff et al., 2004; Hu et al., 2017). Another nsp9 structure was determined featuring a rhinoviral 3C protease sequence LEVLFGQP in the N-terminal tag (PDB 6WQQ). The inserted peptide folds back and forms a β-sheet with the N-terminal of the protein. Residues LEVLF of the peptide make hydrophobic contacts with residues P6, V7, A8, L9, Y31, M101, S105, and L106 of nsp9, and is hydrogen-bonded with residues P6, V7, L9 and S105. These nsp9 residues are conserved among other nsp9 homologs, including full conservation in SARS-CoV. This provides evidence that this binding site could be targeted by a peptidomimetic small molecule to disrupt dimer formation and thus reduce RNA binding and viral replication.

Using FTMap, two potential druggable sites were identified in nsp9. One site is defined by residues R39-V41, F56-S59, I65-E68, I91, and K92, the other one by residues S13, C14, D26-L29, L45-L48, and K86 (Fig. S7B). These two sites lie in regions of intermediate-to-high cryptic site score, and do not overlap with the dimerization interface, but are close to positively charged residues that might be involved in RNA binding, as was postulated for IBV (Chen et al., 2009a); it should also be considered that they might interfere in nsp9-nsp8 PPI, though this is a subject to validation. A hot-spot within a region of moderate-high cryptic site score was identified in the C-terminus of nsp9, delimited by residues C73, F75, L88, L103, L106, A107, and L112 (Fig. S7C); a molecule binding in this region might clash with the N-terminal part of the other protomer.

4.1.10. ADP-ribose-phosphatase (nsp3 domain)

The first nsp3 macrodomain (Mac 1) is conserved throughout CoVs and has an ADP-ribose phosphate (ADRP) activity, by which ADRP binds to and hydrolyzes mono-ADP-ribosylated (Alhammad et al., 2020). This appears to be related to the removal of ADP-ribose from ADP-ribosylated proteins or nucleic acids (RNA and DNA); it should be pointed out that ADRP is able to remove mono-ADP-ribose, but not poly-ADP-ribose (Alhammad et al., 2020). Anti-viral ADP-ribosylation is a host post-translational modification in response to viral infections, since many of the IFN and cytokine signaling components, such as NF-κB essential modulator (NEMO), TANK-binding kinase 1 (TBK1), and NFκB need to be ribosylated to be fully active (Fehr et al., 2020). Although ADRP is not an essential protein for viral replication, it has been shown to be an essential pathogenesis factor in animal models for CoV infection; for example, mutations of ADRP in SARS-CoV enhanced IFN response and reduced viral loads in vivo in mice models (Fehr et al., 2016). Thus, its role against host-induced anti-viral activity makes it an attractive target for drug design.

Recently, five crystal structures of SARS-CoV-2 ADRP were solved (see Table S1), including the apo form (PDBs 6VXS at resolution 2.0 Å, 6WEN at 1.35 Å), and in complex with 2-N-morpholinoethanesulfonic acid (MES) (PDB 6WCF at 1.09 Å), AMP (PDB 6W6Y at 1.45 Å), and ADP-ribose (PDB 6W02 at 1.50 Å) (Fig. S8). Another structure of ADRP complexed with ADP-ribose (PDB 6W0J) at 2.2 Å is available. These structures exhibit low main chain atom RMSD between any pair of them, with values in the range 0.25–0.65 Å. The SARS-CoV-2 ADP-bound ADRP structures share structural similarity to related homologs from SARS-CoV (71% sequence identity, 82% similarity) and MERS-CoV (40% sequence identity, 61% similarity), with main chain RMSD values of 0.6 Å (PDB 2FAV) and 1.4 Å (PDB 5HOL), respectively.

The binding site of SARS-CoV-2 ADRP (Fig. S8) also bears high similarity with those of SARS-CoV and MERS-CoV. The ADP-ribose is stabilized within the binding site through hydrophobic interactions, and direct and solvent-mediated hydrogen-bond interactions; it should be highlighted that most of the hydrophobic contacts involve the main chain atoms. The binding mode of ADP-ribose is conserved in those three CoVs (RMS values of 0.3 Å and 1.2 Å, respectively, with respect to SARS-CoV-2 ADRP), with ADP-ribose exhibiting similar affinities towards the three CoVs (Alhammad et al., 2020). Within the overall conserved binding site conformation of the SARS-CoV-2 ADRP structures, some shifts are observed when comparing the apo structure, and those in complex with AMP, ADP-ribose, and MES, mainly around the proximal ribose. The rotamer states of several side chains (F132, I131, F156) adopt a
ligand-dependent conformation, and a flip in the A129-G130 peptide bond could be observed, dependent on the presence of the phosphate group in ADP-ribose (phosphate 2), or MES.

Considering its highly conserved structural features in CoVs, especially SARS-CoV and MERS-CoV, and its role in countering host-induced antiviral responses, ADRP appears as an attractive therapeutic target. It should be noted, however, that no other druggable binding site could be identified other than the ADP-ribose pocket using FTMap or ICM Pocket Finder, nor could cryptic sites be found on the surface.

4.1.11. Ubiquitin-like 1 domain (nsp3)

The first ~110 residues of nsp3 have an ubiquitin-like fold, and are thus named the Ubl1 domain. The function of Ubl1 in CoVs is related to ssRNA binding, while probably interacting with the N protein too. In SARS-CoV Ubl1 has been shown to bind ssRNA with AUA patterns, and since the SARS-CoV S'-UTR (un-translated region) is rich in AUA repeats, it probably binds to it (Serrano et al., 2007); in fact, SARS-CoV-2 has 439 AUA within its genome. In MHV, this Ubl1 domain binds the structural N protein (Hurst et al., 2013). It was also shown that MHV-Ubl1 is essential for viral replication, since viable virae could not be recovered from the Ubl1 full deletion mutant (Hurst et al., 2013).

There is yet no available experimentally solved structure of SARS-CoV-2 Ubl1. Considering that the sequence identity and similarity with its SARS-CoV counterpart is 79% and 93%, respectively (with only one deletion in the alignment, close to the N-terminal), a structural model was built using SARS-CoV Ubl1 (PDB 2GRI) as template (Fig. S9A). There are several positive residues on the protein surface, compatible with ssRNA binding. These residues are conserved in SARS-CoV, with the exception of R23N, R102H, and N98K (SARS-CoV numbering). Two small potentially druggable sites were identified with FTMap and ICM Pocket Finder, which lie in areas of above-average cryptic site score. One site is defined by the side chains of Y42, T43, T48, E52, F53, C55 and V56 (Fig. S9B), and the second one by F25-D28, T86, Y87, W82, and C104-F106 (Fig. S9C). These sites are near positively charged residues, but do not overlap with them. While a molecule binding to them might interfere with ssRNA binding, it is also possible that it might disrupt PPIs with partner proteins, such as the N protein. Further biochemical and functional characterization of Ubl1 is needed to shed light on the actual value of these sites.

4.1.12. SARS-unique domain (SUD) (nsp3 domain)

The SUDs is part of nsp3 and it is implicated in tracking the viral RNA to DMVs. In SARS-CoV-2 and SARS-CoV there are three SUD domains connected by short peptide linkers: SUD-N, SUD-M, and SUD-C, indicating the N-terminal, the middle, and the C-terminal regions of SUD, respectively. SUD-N (macromolecular domain 2, Mac 2) binds G-quadruplexes, an unusual nucleic-acid structures formed by guanine-rich nucleotides in ssRNAs; it shares 71% of identity and 85% similarity with its SARS-CoV counterpart. SUD-M/Mac3 (macromolecular domain 3, Mac3) has a single-stranded purine rich (G and/or A) RNA binding activity (Johnson et al., 2010) which includes poly(A) and G-quadruplexes; this allows SUD-M to act as a poly(A)-binding-protein and thus protect the poly(A) tail at the 3’ end of viral RNA from host 3’ exonucleases. SUD-C/DPUP (Domain Preceding Ubl2 and PlP10) also binds to ssRNA, and recognizes purine bases more strongly than pyrimidine bases (Johnson et al., 2010). This RNA binding activity is apparently stabilized by the presence of SUD-M, while SUD-C seems to modulate the secondary structure of SUD-M (Johnson et al., 2010). Mutagenesis analysis in SARS-CoV showed that SUD-M is indispensable for virus replication, while the absence of SUD-N or SUD-C barely reduce the virus titer (Kusov et al., 2015).

The structure of the SARS-CoV construct SUD-N/SUD-M (SUD-NM) has been solved by crystallography at 2.2 Å resolution (PDB 2W2G). The solution structures of SARS-CoV SUD-M and SUD-C within a SUD-MC construct were obtained using NMR (PDBs 2KQV and 2KQW, respectively), together with the isolated SUD-C (PDB 2KA1). The isolated SARS-CoV SUD-M has also been solved by NMR (PDBs 2JZD and 2JZE), together with the structure of an N-terminal extended SUD-M (PDBs 2RNK and 2JZD). Main chain RMD values among corresponding solved structures are within 0.8 Å. It has been shown that SUD-NM is monomeric in solution (Johnson et al., 2010), and the absence of evidence suggesting a tight transient or static contacts in solution showed that SUD could be modeled as three flexible linked globular domains (Johnson et al., 2010).

The SARS-CoV-2 SUD domains are very similar to their SARS-CoV counterpart, with sequence identity (similarity) of 71% (91%), 79% (94%), and 72% (92%) for SUD-N, SUD-M, and SUD-C, respectively. Considering that only SUD-M appears to be essential to viral replication, we focused our druggability analysis on it. We thus built a model by homology using PDB 2AKF as template. For SARS-CoV, it was shown that upon poly(A) binding to SUD-M, the molecular surface area of SUD-M affected by NMR chemical shift perturbation experiments was mapped to a positively charged surface cavity (Johnson et al., 2010), defined by residues N532, I533, I556, M557, A558, T559, Q561, and V611 (Chatterjee et al., 2009). This area is wholly conserved in SARS-CoV-2, and a potential druggable binding site within it was identified using FTMap and ICM Pocket Finder (Fig. S10). It could be hypothesized that a small-molecule binding to this site might preclude RNA binding.

4.1.13. Nucleic-acid binding region (NAB, nsp3 domain)

NAB is a small domain of ~120 amino acids that binds ssRNA, strongly preferring sequences containing repeats of three consecutive guanosines (Serrano et al., 2009). In SARS-CoV, NAB binds ssRNA through a positively charged surface patch defined by the residues K75, K76, K99, and R106, while the neighboring residues N17, A18, S19, D66, H69, T97 are also affected by RNA binding (Serrano et al., 2009). Interestingly, this RNA binding site bears similarity to that of the sterile alpha motif of the Saccharomyces cerevisiae Vts1p protein (Serrano et al., 2009). Experiments have shown that N-terminal and C-terminal extensions, corresponding to links with the PlP10 and the TM-Lumen/Ectodomain, respectively, behave as flexibly disordered segments (Serrano et al., 2009).

While the SARS-CoV-2 NAB structure is not yet available, the SARS-CoV counterpart has been solved by NMR (PDB 2KB7). We used the latter to build a structural model of SARS-CoV-2 NAB by homology (sequence identity 81%, similarity 94%, no gaps in the alignment). All the positive residues are conserved, with two additions, T43K and S51K (Fig. S11A).

No druggable binding sites were identified within the positive charged patch on the surface where ssRNA binds in SARS-CoV; however, two cryptic sites with nearby CSs were predicted on the sides of that patch (Fig. S11B). It should be further explored whether these sites are involved in PPI, and whether small-molecules binding to them might allosterically modulate ssRNA binding.

4.1.14. Other nsp3 domains

The DUF3655/HVR/Acidic-domain, TM-Lumen/Ectodomain, and the C-Terminal domain/Y-Domain (Table S1) are structurally characterized nsp3 domains, which bear little similarity to any other experimentally solved structure, thus ruling out the possibility of homology modeling.

DUF3655-HVR is a hypervariable, Glu and Asp rich domain, probably implicated in protein-protein interactions with N, as in MHV (Keune and Giedroc, 2013). Since it is not essential to viral replication in MHV (Hurst et al., 2013), it would be a second priority for drug targeting.

The TM-Lumen/Ectodomain is flanked by two transmembrane domains (TMs), exposed to the ER lumen, and works by binding to the lumen domain of nsp4, necessary to form the DMVs where CoVs replicate, anchoring the whole nsp3 protein to membranes through the TMs. While DMVs formation is essential, the lack of structural information
poses an insurmountable hurdle for drug discovery (Hagemeijer et al., 2014).

The C-Terminal domain/Y-Domain is conserved in CoVs, and it seems to be involved in the DMVs formation, probably interacting with nsp6, and improving nsp3-nsp4 interaction (Hagemeijer et al., 2014). Although breaking this interaction would seriously impact viral replication, the lack of structural and biochemical information precludes any targeting attempt.

4.1.15. Other non-structural proteins: nsp1, nsp2, nsp4, and nsp6

The nsp1 is the leader protein and the first translated and PL pro-processed protein. It has the capacity to bind to the 40S ribosomal subunit to inactivate the translation of host mRNAs (Thoms et al., 2020), also selectively promoting host mRNAs degradation, which makes it the main actor in host cell shutdown (Thoms et al., 2020), and indirectly in the immune response evasion by delaying IFN responses (Lei et al., 2020). Additionally, based on recent proteomics analyses, SARS-CoV-2 nsp1 seems to interfere with the host DNA duplication by interacting with the DNAPoA complex, which is the host primase complex, and the first step in DNA synthesis (Gordon et al., 2020b). Although nsp1 is not essential for viral replication, its absence makes the virus susceptible to IFN (Lei et al., 2020), which would make it an important pathogenic factor and a good target for drug design. However, the structural information available of this 180-amino acids protein is limited. An NMR structure from SARS-CoV (PDB 2GD7) of 116 amino acids (corresponding to H13 to G127 in SARS-CoV-2) does not contain the 40S binding region (which includes residues K164 and H165), and only partially represents the active site for RNA binding (Lokugamage et al., 2012). There is a recent cryo-EM structure of the nsp1 C-terminus (E148-G180) in complex with the 40S ribosomal subunit and RNA (PDB 6ZLW). While amino-acids K164 and H165 are present, the C-terminal portion of only 33 residues cannot be used for structure-based drug design.

The nsp2 is a membrane protein not essential for the viral production in SARS-CoV and MHV homologs. However, in MHV, the presence of nsp2 positively affects the viral titer (Gadlage et al., 2008). It is a delocalized membrane protein, but is recruited to replication sites by other viral components (Graham et al., 2005), probably by the M protein (Li et al., 2020a). In SARS-CoV-2, proteomic data showed that nsp2 interacts with cellular components associated with vesicles formation and translational regulators (Gordon et al., 2020b), which might hint that its function is associated with the formation of membrane structures and co-opting host components. However, up to the present date there is no structural data of nsp2 from SARS-CoV-2 or related proteins.

The nsp4 is essential for viral replication in MHC (Sparks et al., 2007). The main function of nsp4 is the formation of DMVs in SARS-CoV and MERS-CoV (Angelini et al., 2013). Nsp4 has four TM domains, a ER-Lumen exposed domain, and a cytoplasmatic exposed C-Terminal domain (Clementz et al., 2008). In SARS and MHV, the lumen domain of nsp4 interacts strongly with the nsp3 lumen domain/ectodomain to induce DMVs formation (Hagemeijer et al., 2014). Thus, an nsp3-nsp4 structure might help to design a PIPI inhibitor that could prevent DMVs formation. The cytoplasmatic C-Terminal domain (~90 amino acids) of MHV and Feline CoV nsp4 structures are available (3VC8, 3GZI, respectively), but SARS-CoV-2 nsp4 shares with them only 59% and 39% sequence identity, respectively.

The nsp6 is a membrane protein with a medium conservation degree between different CoVs. In SARS-CoV nsp6 collaborates with nsp4 and nsp3 in DMVs formation, also inducing the formation of membrane vesicles (Angelini et al., 2013). Additionally, in several CoVs, nsp6 activates omegasome and autophagosome formation independently of starvation, to degrade cellular components and so increase the availability of resources for viral replication (Cottam et al., 2011). Nevertheless, there is no structural information on any related protein which would allow homology modeling and structure-based drug discovery.

4.2. Structural proteins

4.2.1. S protein

The SARS-CoV-2 spike-protein (S protein, ORF2) is a large homotrimeric multidomain glycoprotein. The small C-terminal transmembrane attaching domain is followed by the S1 and S2 subunits. During infection, the receptor binding domain (RBD) in S1 is exposed and recognized by ACE2, the junction S1/S2 is cleaved by a furin-like protease releasing the S1 domain, while S2 is cleaved again by the metallloprotease TMPRSS2, to expose the FP, which is responsible in inducing the membrane fusion mechanisms (Wrapp et al., 2020) (Fig. 1).

Several strategies are focused on inhibiting the viral entry, either by interfering with the binding of S to ACE2, or with the membrane fusion induction (Zhou et al., 2020). The first approach was the development of inhibitory antibodies or recombinant ACE2 proteins that block the S protein (Zhou et al., 2020); as the outer surface component of CoVs, the S protein is a major target of antibodies, and the main focus of vaccine development. Using a different approach, based on structural information and the knowledge of the membrane fusion mechanisms, several peptidomimetics have been developed, such as a series of lipo-peptides EK1C1-EK1C7 and IPB02, that target the subdomain HR1 in the S2 fragment to inhibit membrane fusion by interfering with the FP, and which showed a decrease of SARS-CoV-2 virus titer in cell cultures (Zhu et al., 2020).

As the mechanism of the virus entry depends on host factors, targeting ACE2 and TMPRSS2 is being explored. There are no commercial ACE2-inhibitors. The main candidates for inhibiting the ACE2-Spike interaction appear to be specific antibodies, which are currently raising great expectations, and are completing their clinical phases. The inhibition of TMPRSS2 is also being explored, by blocking the proteolytic priming of the S2 fragment, necessary to induce membrane fusion; the inhibitors Nafamostat, Gabexate, and Calmostat (Shrimp et al., 2020) have been shown to inhibit viral production in cell cultures, and will be tested in humans; however, their low half-life suggests a need for developing more efficient new drugs [cf. Ref (Singh et al., 2020), for new options targeting TMPRSS2].

4.2.2. E protein

It is a small homopentameric membrane protein (75 amino-acids per protomer) (Pervushin et al., 2009), which has been shown to be essential for viral particle assembly in SARS-CoV (Siu et al., 2008). The N-terminal region of the protein spans the lipid bilayer twice, while the C-terminal is exposed to the interior of the virus (Shen et al., 2003). In IBV and SARS-CoV, the E protein interacts with the viral M protein through an undefined region (Chen et al., 2009b); and also through its C-terminus with the host Protein Associated with Lin Seven 1 (PALS1), a factor associated with pathogenesis (Teoh et al., 2010). In many CoVs, the E protein works as an ion-channeling viroporin (Pervushin et al., 2009), which affects the production of cytokines, and consequently the inflammatory response (Nieto-Torres et al., 2014).

The NMR structures of the SARS-CoV-2 and SARS-CoV E protein homopentamers (PDB 7K3G and 5X29, respectively) are shown in Fig. S12. Only the TM regions were solved, showing 31 and 58 residues per helix, respectively. While in SARS-CoV-2 the helices are parallel, and in SARS-CoV they exhibit a tilt angle of ~15°, the size of the channel is roughly conserved. In fact, based on structural and functional considerations, the E protein channel would be the primary target site for drug development. Moreover, hexamethylene amiloride (HMA) has been reported to bind to the SARS-CoV E protein homopentamer, but not to an isolated protomer (Surya et al., 2018).

4.2.3. N protein

β-coronavirus nucleocapsid (N) proteins are involved in the packing of viral +ssRNA to form a ribonucleoprotein (RNP) complex, which interacts with the M protein (He et al., 2004). They share an overall conserved domain structure, with an RNA-binding N-terminal domain
and association of N protein homotetramers (Ye et al., 2020). Since the formation of the RNP complex is essential for viral replication, identification of small-molecule modulators of the nucleocapsid assembly, interfering with NTD RNA binding, or precluding CTD dimerization or oligomerization, would be valid therapeutic strategies against SARS-CoV-2.

Several structures of the N protein NTD and CTD have recently become available (see Table S5). Structures of the NTD (and CTD) superimpose closely among themselves. With respect to SARS-CoV, the NTD and CTD share 88% (96%), and 96% (98%) sequence identity (similarity), respectively, and the corresponding structures overlay with RMSD values within 0.8 Å.

The NTD structure solved by NMR (PDB 6Y13) showing that amino-acids A50, T57, H59, R92, I94, S105, R107, R149, and Y172 participate in RNA binding (Dinesh et al., 2020) is shown in Fig. S13A. A druggable site identified near those residues partially overlaps with the AMP binding site in the NTD of the homolog protein human CoV OC43 (HG0-COC43) (PDB 4L4U), in agreement with an earlier hypothesis (Kang et al., 2020). Two small borderline druggable sites have also been identified with FTMaP (site 1: L159-P162, T165, L167, A173, S176; site 2: Q70-N75, Q83, T135, P162). These sites would not overlap with RNA binding, and further experiments are needed to confirm whether they could be allosteric sites, or near PPI interfaces.

The CTD homodimeric structure is shown in Fig. S13B (PDB 6WZQ), also displaying two potential druggable sites. A molecule binding to any of them might interfere with dimer formation. Two other distinct sites were predicted on the surface of the homodimer, one over the central four-stranded β-sheet, and the other opposite to it, near the C-terminal α-helices (Fig. S13C). Since it is not clear yet how the homotetramer is formed (Ye et al., 2020), it should be further explored whether any of these sites might overlap with PPI interfaces.

4.2.4. M protein

It is a membrane homodimeric glycoprotein that forms part of the virion. In SARS-CoV, the M protein interacts with the N protein, being a nexus between virus membrane formation and RNA association in the virion (Nal et al., 2005), also inhibiting IFN production in SARS and MERS-related diseases (Lai et al., 2016; Sui et al., 2014). While it is established that dimerization and N-M PPI motifs reside in the C-terminal α-helices (Fig. S13C), it is still unclear how the homotetramer is formed (Ye et al., 2020), it should be further explored whether any of these sites might overlap with PPI interfaces.

4.3. Accessory proteins

4.3.1. Orf3a/X1/U274

The orf3a is characterized as a potassium ion channel in SARS-CoV, involved in inducing caspase-dependent apoptosis under different pathological conditions (Chan et al., 2009). It is included in the virion (Shen et al., 2005), and interacts specifically with the M, E, and S structural proteins, as well as with orf7a/U122 (Tan et al., 2004). In SARS-CoV, orf3a expression increases the mRNA levels of all three subunits of fibrinogen, thus promoting fibrosis, one of the serious pathogenic aspects of SARS (Tan et al., 2005), and the expression of NFκB, IL8, and JNK, all involved in inflammatory responses (Kanzawa et al., 2006). In this scenario, design therapeutics that suppress its function could be very important; moreover, the presence of similar proteins to orf3a in other β-CoVs (SARS-CoV-2 orf3a shares ~73% identity and ~85% similarity with its SARS-CoV counterpart), and in α-CoVs, suggests that drugs targeting orf3a might be a therapeutic option against a broad range of CoV-related diseases (Kern et al., 2020).

Recently the SARS-CoV-2 Orf3a structure was solved by cryo-EM (PDB 6XDC). The N-terminus (amino acids 1–39), the C-terminus (239–275), and a short loop (175–180) were not observed, probably due to molecular disorder. The orf3a was solved as a homodimer, although the authors were able to reconstruct the tetramer at a lower final resolution of ~6.5 Å, also showing a model of the neighboring dimers, and inferring from this model that residues W131, R134, K136, H150, T151, N152, C153, and D155 were involved in a network of interactions that would mediate tetramerization (Kern et al., 2020) (Fig. 11A). Considering that SARS-CoV orf3a has been identified as an emodin-sensitive potassium-permeable cation channel, the narrow size of the pore in the SARS-CoV-2 orf3a structure strongly suggests that it is in the closed or inactive conformation (Kern et al., 2020).

Using FTMaP and ICM Pocket Finder, a druggable site was found within the dimer, lined up by the side chains L65, L71, Y141, D142, N144, P159, N161, and Y189 of both protomers (Fig. 11B). This cavity has also been identified in Ref. (Kern et al., 2020). Since conformational changes of the TMs are needed for channel opening, we could hypothesize that a small-molecule binding within this site would interfere with this rearrangement, or directly block the channel. Another potentially druggable site delimited by residues K132-R134, C148-H150, D155, Y156, C200-V202, and H204, was identified, which partially overlaps with the proposed tetramerization interface, and thus it could be explored for a possible PPI inhibitor (Fig. 11C).

4.3.2. Orf7a/X4/U122

The SARS-CoV-2 orf7a is a transmembrane protein of 121 amino acids (106 if only the mature protein is considered, excluding the signaling peptide), with 86% identity and 94% similarity with respect to its SARS-CoV counterpart. Interestingly, it exhibits structural similarity to Igs (Nelson et al., 2005; Tan et al., 2020), but shares poor sequence identity with proteins of the Ig superfamily (within the 2–16% range). Like orf3a, orf7a is also included in the virion, and both proteins have been shown to interact with each other in SARS-CoV (Tan et al., 2004) and in SARS-CoV-2 (Li et al., 2020a). In SARS-CoV, orf7a expression increases the expression of NFκB, IL8 and JNK (Kanzawa et al., 2006), while the deletion of orf7a reduces the virus titer 30-fold. Orf7a appears to be unique to SARS CoVs, showing no significant similarity to any other protein, either viral or non-viral (Nelson et al., 2005). In SARS-CoV-2, orf7a interacts with midasin AAA ATPase 1 (MDN1) and HEAT repeat containing 3 (HEATR3) (Gordon et al., 2020b). MDN1 is a protein involved in the maturation of ribosomes in eucaryotes (Garbarino and Gibbons, 2002), while HEATR3 has a positive role in Nucleotide-binding oligomerization domain-containing protein 2 (NOD2)-mediated NF-κB signaling (Zhang et al., 2013), and is also involved in the assembly of the SS ribosomal subunit (Calvino et al., 2015). From this information it can be expected that orf7a works by regulating inflammation like its SARS-CoV homolog, and also that it is involved in protein translation regulation. The N-terminal ectodomain structure of orf7a (amino acids 1–66, using the numbering of the mature protein) is available for SARS-CoV, either from crystallography (PDB 1XAK at 1.8 Å) or NMR (PDB 1OY4), and recently for SARS-CoV-2 (PDB 6W37). The structure displays a seven-stranded β-sandwich fold with two β-strand packed loops (Fig. S14), and exhibits the main chain RMSD values of 0.4 Å and 0.9 Å with the corresponding SARS-CoV orf7a structures, respectively. The SARS-CoV NMR structure also features a disordered part corresponding to the stalk region, between the ectodomain and the membrane (residues 68–82). The transmembrane region comprises approximately amino acids 83–101.

While the functionality and interaction partners of orf7a are not clearly known, we identified two hot-spots on the surface of orf7a. An extended cryptic site was predicted lined up by residues H4, Q6, L16,
P19, Y60, L62, with a CS nearby which includes a deep hydrophobic pocket already identified in SARS-CoV as a potential PPI site (Nelson et al., 2005) (Fig. S14). A hot-spot defined by residues E18, T24, Y25, F31, P33, A35, N37, and F50 was also identified (Fig. S14). Both sites might have functional roles, and further studies are needed to confirm this hypothesis, and whether small-molecules binding to any of them would interfere with the viral mechanism.

4.3.3. Orf8

Orf8 is one of the fastest evolving genes in SARS-CoV-2, as can be inferred by its high variability (Tan et al., 2020). The orf8 expression is not essential for SARS-CoV and SARS-CoV-2 replication; it was observed, however, that a 29 nucleotide deletion in SARS-CoV may correlate with milder disease (Muth et al., 2018). SARS-CoV-2 orf8 is involved in inhibiting IFN signaling (Li et al., 2020b) and also downregulates microtubule organization and IRES dependent translation factors (Gordon et al., 2020).

The SARS-CoV-2 orf8 shares less than 20% sequence identity with its SARS-CoV counterpart, which might be taken into consideration to understand the lethal characteristics of SARS-CoV-2. The crystal structure of SARS-CoV-2 orf8 solved at 2.04 Å exhibits an Ig-like fold, and revealed two novel dimerization interfaces unique to SARS-CoV-2: i) a covalent dimer-interface formed through a sequence-specific region, and linked by an intermolecular disulfide-bond mediated by the unique C20 of each monomer, and ii) a non-covalent interface stabilized by the unique I76 motif defined by residues I47-L60, V62, D63, P19, Y60, L62, with a CS nearby which includes a deep hydrophobic pocket already identified in SARS-CoV as a potential PPI site (Nelson et al., 2005) (Fig. S14). A hot-spot defined by residues E18, T24, Y25, F31, P33, A35, N37, and F50 was also identified (Fig. S14). Both sites might have functional roles, and further studies are needed to confirm this hypothesis, and whether small-molecules binding to any of them would interfere with the viral mechanism.

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essential for replication, they have a lower natural selection pressure, and therefore a higher mutation rate. In fact, the deletion of orf3a, orf3b, orf6, orf7a, and orf7b in SARS-CoV do not eliminate viral production in vitro (Yount et al., 2005), nevertheless, it does affect pathogenicity and virulence.

The following proteins lack experimentally solved structure, or they cannot be modeled due to the absence of similar experimentally solved proteins which could serve as templates:

Orf3b is a short protein observed in some β- and γ-coronaviruses, and produced by an alternative start-codon. In SARS-CoV it was shown to inhibit the expression of IFN-β during synthesis and signaling (Kopecky-Bromberg et al., 2007). It differs considerably among different CoVs, but maintains its pathogenic function (Zhou et al., 2012).

The SARS-CoV orf6 is a small membrane protein (Tangudu et al., 2007) sharing 69% of identity with its SARS-CoV-2 counterpart. It acts as a pathogenicity factor, due to its ability to convert a sublethal MHV infection into a lethal one (Pewe et al., 2005); this property only depends on the N-terminal transmembrane segment (Netland et al., 2007). The SARS-CoV orf6 enhances viral replication (Zhao et al., 2009) through interaction with components of the viral replication machinery, such as nsp8 (Kumar et al., 2007). In addition, it shows pathogenic activity due to its ability to induce apoptosis, similar to orf3a and orf7a. Additionally, SARS-CoV-2 orf6 is involved in the inhibition of IFN signaling, as its SARS-CoV paralog (Li et al., 2020b).

In SARS-CoV, orf7b is a small membrane protein that could be included in the virion (Schaecher et al., 2007), and is probably involved in attenuating viral production (Pfefferle et al., 2009).

Orf9c/orf14 is the third putative protein translated from the orf 9 (Shukla and Hilgenfeld, 2015), and the least characterized. It enables immune evasion and coordinates cellular changes, impairing IFN signaling, antigen presentation, and complement signaling, while inducing interleukin-6 (IL-6) signaling (Dominguez Andres et al., 2020).

Orf10 is a putative membrane protein present only in SARS related CoVs (Caglani et al., 2020). No functional activity has been determined experimentally, but it is inferred from proteomic data that it is involved in targeting host proteins for proteasome degradation (Gordon et al., 2020b).

5. SARS-CoV-2 polymorphisms

Understanding the evolution of pathogens is important for the effective development of new antiviral strategies to circumvent drug resistance.

Since the outbreak in December 2019, several genetic variants have appeared compared to the first SARS-CoV-2 genome sequenced in Wuhan (Mercatelli and Giorgi, 2020). These polymorphic amino acids are found in structural, non-structural, and accessory proteins, and the most frequent ones are listed in Table S6. In structural proteins, the most notorious case is the D614G mutation in the RBD domain of the S protein, which became predominant during the first months of the outbreak (Plante et al., 2020), along with other changes such as the D936Y in the FP domain (Cavallo and Oliva, 2020). In non-structural proteins, the P504L and Y541C are the predominant mutations in the helicase nsp13 (Table S6), and they were reported to affect the RNA binding (Begum et al., 2020). Among accessory proteins, the mutation Q57H in orf3a is becoming dominant and appears to be associated with disease severity; orf3a also possesses other relevant polymorphism such as S171L, G196V and G251V (Issa et al., 2020) (Table S6).

Knowledge of polymorphic amino-acids could be taken into consideration when selecting molecular targets for drug development to avoid or reduce potential drug resistance.

6. Discussion and perspective

The appearance of COVID-19 caused by the SARS-CoV-2, its fast spread throughout the world, and the mounting number of infected persons have triggered a prompt and resolute quest for therapeutic options to treat this serious infectious disease.

In a recent work, the SARS-CoV-2-human interactome was characterized, and >300 PPIs were identified. It was found that 66 of the interacting host factors could be modulated with 69 compounds, including some approved drugs (Gordon et al., 2020b). This study will certainly enhance the possibilities of drug discovery efforts targeting host cells receptors. As of today, candidates include the antimalarial chloroquine and its derivatives (Zou et al., 2020), anti-inflammatory drugs and immunomodulators, and even anticoagulants and anti-fibrinolytic drugs (Hoffmann et al., 2020).

Although the factors that condition the progression of the disease in severe cases are not fully understood, evidence seems to show that the symptoms in patients with moderate to severe disease due to SARS-CoV-2 are related not only to viral proliferation, but also to two factors related to its pathogenesis: i) an exacerbated inflammatory response associated with increased concentrations of proinflammatory cytokines, such as tumor necrosis factor-α (TNF-α) and ILs, including IL-1 and IL-6 (Hadjadj et al., 2020; Mehta et al., 2020), and ii) abnormalities in coagulation and thrombosis, similar to a combination of mild disseminated intravascular coagulopathy and a localized pulmonary thrombotic microangiopathy, which could have a substantial impact on organ dysfunction in the more severely affected patients (Levi et al., 2020).

It is clear that a thorough characterization of the druggability of the SARS-CoV-2 proteome would provide a rich array of alternative targets for drug discovery. In this work we present an in-depth functional, structural and druggability assessment of all non-structural, structural, and accessory proteins of SARS-CoV-2, identifying potential druggable allosteric and PPI sites throughout the whole proteome, thus broadening the repertoire of current targetable proteins. It should be stressed that druggability characterization of a site does not necessarily imply that a compound binding at that site will modulate that target and exhibit an observable biological effect.

We would like to highlight three interesting drug discovery strategies:

- The helicase (nsp13), and methyltransferases N7-Mtase (nsp14) and 2′-O-MTase (nsp16) contribute to genome stability through their involvement in the capping process; interfering with the functioning of one or more of these proteins could be an excellent antiviral approach.
- The ExoN (nsp14) is responsible for proofreading, and thus for the extremely low mutation rate and nucleoside analogs resistance of SARS-CoV-2; a therapeutic combo of Remdesivir (or other RdRp nucleoside compound) in combination with an ExoN inhibitor could be a valid approach to boost the efficacy of nucleoside-based therapies.
- In CoVs, a delayed IFN response is a redundant pattern that allows robust viral replication, and also induces the accumulation of cytokine-producing macrophages, thus increasing the severity of the disease (Chen and Subbarao, 2007). Targeting one (or a combination) of NendoU (nsp15), ADRP, PLP1, orf6, and orf9b, would certainly obstruct viral replication.

We are convinced that our work will contribute to the quick development of an effective SARS-CoV-2 antiviral strategy which, in view of the high similarity among CoVs, might be useful to fight related viruses. Moreover, these therapeutic options might be instrumental in fighting different CoV-associated diseases that could threaten global health in the future.

Competing interests

The Authors declare that no competing interests exist.
CRediT authorship contribution statement

Claudio N. Cavasotto: conceived the original idea of the work, designed the research process, performed the functional, structural and druggability, Formal analysis, interpreted the results, wrote the manuscript. Maximiliano Sánchez Lamas: contributed to the, Formal analysis, interpretation of results, and to the manuscript. Julián Maggini: Formal analysis, interpretation of results, and to the manuscript. All authors have reviewed the manuscript and approved the submission.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ejphar.2020.173705.

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