Capping of viral messenger RNAs is essential for efficient translation, for virus replication, and for preventing detection by the host cell innate response system. The SARS-CoV-2 genome encodes the 2'-O-methyltransferase nsp16, which, when bound to the coactivator nsp10, uses S-adenosylmethionine (SAM) as a donor to transfer a methyl group to the first ribonucleotide of the mRNA in the final step of viral mRNA capping. Here, we provide biochemical and structural evidence that this reaction requires divalent cations, preferably Mn2+, and a coronavirus-specific four-residue insert. We determined the x-ray structures of the SARS-CoV-2 2'-O-methyltransferase (the nsp16-nsp10 heterodimer) in complex with its reaction substrates, products, and divalent metal cations. These structural snapshots revealed that metal ions and the insert stabilize interactions between the capped RNA and nsp16, resulting in the precise alignment of the ribonucleotides in the active site. Comparison of available structures of 2'-O-methyltransferases with capped RNAs from different organisms revealed that the four-residue insert unique to coronavirus nsp16 alters the backbone conformation of the capped RNA in the binding groove, thereby promoting catalysis. This insert is highly conserved across coronaviruses, and its absence in mammalian methyltransferases makes this region a promising site for structure-guided drug design of selective coronavirus inhibitors.

INTRODUCTION

The recently emerged human pathogen severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is a positive-stranded RNA virus responsible for the ongoing pandemic of highly transmissible fatal respiratory coronavirus infectious disease (COVID-19), which has caused more than 3 million deaths worldwide (1). Like other coronaviruses, SARS-CoV-2 carries a large RNA genome of ~30 kilobases (2), which is directly translated into two polyproteins, Orf1a and Orf1ab. The polyproteins are subsequently autoprocessed into the nonstructural proteins (nspS) that comprise the replication and transcription complex (RTC). The genome also serves as a template for viral RNA transcription to generate nine canonical subgenomic mRNAs that encode the SARS-CoV-2 structural and accessory proteins (3). These mRNAs are generated by the RTC through a discontinuous transcription process and thus have identical 5'-ends with an adenosine ribonucleotide at the first position (A1) (4, 5). These mRNAs are further modified by 5'-capping to stabilize, improve translation, and protect the RNA from surveillance by the host innate immune system (6, 7).

Coronavirus genome replication and transcription processes are confined to viral-specific organelles formed from endoplasmic reticulum membranes (8, 9) during early-stage infection, and probably mitochondrial, endosomal, and Golgi membranes during viral particle assembly (8–10). Thus, nascent coronavirus mRNAs do not have access to the host mRNA capping and methylation machinery in the nucleus but instead encode their own capping machinery.

The viral replication helicase nsp13 also has 5'-RNA triphosphatase activity that removes the 5'-γ-phosphate from the nascent ppp-RNA (11). The RNA-dependent RNA-polymerase nsp12 can then transfer a guanosine monophosphate to the 5'-end of the mRNA (12, 13), and this guanosine is then methylated by the nsp14-nsp10 heterodimer to generate N7-methylated Cap-0-RNA (m7GpppA1-RNA). Last, to form the Cap-1-RNA, a methyl group is transferred from S-adenosylmethionine (SAM) to the 2'-OH of the first adenosine ribonucleotide. For coronaviruses, this last reaction is catalyzed by the 2'-O-methyltransferase (MTase), a heterodimeric complex of nsp16 with the activator nsp10 (Fig. 1A) (11, 14, 15). The coronavirus 2'-O-MTase is a validated antiviral drug target, because the formation of the nsp16-nsp10 complex can be blocked by peptides and protect mice from lethal challenge with mouse hepatitis virus (MHV) (16, 17).

Biochemical studies of the RNA MTases from SARS-CoV and Middle East respiratory syndrome coronavirus (MERS-CoV) demonstrated that methyl transfer is activated by divalent cations. SARS-CoV nsp14 is activated by Mn2+, but not by Mg2+ (11), whereas nsp16 can be activated by Mn5+ or Mg2+ and to a lesser extent by Ca2+ (11, 18–20). These data are consistent with known requirements for divalent cations by other viral RNA MTases, including dengue virus (DENV) NS5, which is stimulated by Mg2+ (21). Furthermore, studies of nsp16 from SARS-CoV and MERS-CoV have demonstrated that the methyl transfer reaction is most efficient in vitro with an RNA substrate composed of at least five ribonucleotides (11, 19, 20, 22). However, the role of metals in nsp16 catalysis and the influence of mRNA length on the reaction are not clear, particularly because no structures have been solved for any coronavirus nsp16 enzyme with bound metals or RNA. To support drug discovery efforts targeting nsp16, researchers around the globe have determined crystal structures of SARS-CoV-2 nsp16-nsp10 in complex with a variety of ligands (23–29), complementing prior structural biology observations for this enzyme from SARS-CoV and MERS-CoV (19, 20, 30).
Fig. 1. Metal ions catalyze 2′-O-methyl transfer and orient the Cap-0-RNA in the active site. (A) Schematic representation of the 2′-O-methyl transfer reaction generating Cap-1-RNA from Cap-0-RNA. The methylated guanosine cap (m7G) moiety is colored in blue, and the methyl group added to the A1 2′-OH position from the methyl donor SAM is in red. The remainder of the RNA structure with the first three labeled ribonucleotides is in black. (B) MTase-Glo luminescence assay results for Cap-0-RNA (m7GpppAUUAAA) and Cap-0 analog (m7GpppA) as substrates. n.d., no activity detected. The nsp16-nsp10 activity with Cap-0-RNA, SAM, and Mg2+ was selected as a reference (100%), and all measured activities were normalized to this value. All values are means ± SD for biological triplicates conducted in two separate experiments using two independent preparations of nsp16-nsp10 (n = 6). (C) The overall view of nsp16-nsp10 in complex with Mg2+, Cap-0-RNA, and SAM (PDB 7JYY). The high-affinity binding site (HBS) and the low-affinity RNA binding (LBS) are labeled. (D and E) Close-up views of nsp16-nsp10 in complex with (D) Mg2+, Cap-0-RNA, and SAM (PDB 7JYY) or with (D) Mn2+, Cap-1-RNA, and SAH (PDB 7L6R). The nsp16 and nsp10 proteins are represented as solvent-exposed surfaces in tan and teal, respectively. Capped RNAs, SAM, and SAH are shown as sticks. Carbons are in gray for capped RNAs and in green for SAM and SAH; oxygens are in red; nitrogens are in blue; phosphates are in orange; sulfur is in yellow. Metal ions are shown as large spheres colored in purple for Mg2+ and orange for Mn2+. Water molecules are small, cyan spheres. Hydrogen bonds between metal ions and waters from the first hydration sphere are shown as black dashed lines.

This information has facilitated detailed examination of the SARS-CoV-2 2′-O-MTase and revealed conformational flexibility of loop 1 and loop8-n3-loop9, loops that border the Cap-0 binding site (25, 26). Despite this extensive structural information, a major gap exists in the understanding of the role of metal ions in the nsp16 2′-O-methyl transfer reaction and the position of ribonucleotides in the RNA binding groove because structures of the enzyme in complex with RNA and metals have not been reported. Herein, we present structures of the SARS-CoV-2 2′-O-MTase nsp16-nsp10 heterodimer in complex with substrates and products in the presence of divalent cations. Our structures reveal important information regarding interactions of nsp16 residues with the capped RNA, the influence of a coronavirus MTase-specific four-residue insert on the conformation of capped RNA, and the role of metal ions in 2′-O-methyl transfer.

RESULTS

SARS-CoV-2 nsp16 activity requires extended capped RNA and divalent cations

We launched this research by confirming that the SARS-CoV-2 2′-O-MTase activity requires divalent cations. We used a custom-synthesized Cap-0-RNA substrate composed of the N7-methylated guanosine (m7G) attached through a triphosphate bridge to a short RNA (AUUAAA), which matches the naturally occurring ribonucleotides at the 5′-end of SARS-CoV-2 mRNAs (31). At a concentration of 2 mM, both Mg2+ and Mn2+ increased MTase activity with this substrate (Fig. 1B). In contrast, 2 mM Ca2+ yielded only 50% of the activity observed with Mg2+, and Na+ did not stimulate activity at 3 mM (Fig. 1B). These data are consistent with observations for SARS-CoV nsp16 (11, 19, 20). The activity of SARS-CoV-2 nsp16 with the Cap-0-RNA substrate (m7GpppAUUAAA) was over 10 times higher than that with the Cap-0 analog (m7GpppA) (Fig. 1B).

Isothermal titration calorimetry (ITC) measurements showed that, although it is a poor substrate for catalysis, m7GpppA does bind to nsp16-nsp10 (Kd = 6.6 ± 0.3 μM) with threefold higher affinity than to nsp16 alone (Kd = 28.0 ± 5.5 μM) (fig. S1D and table S1). In contrast, m7GpppG bound only to the nsp16-nsp10 heterodimer but not to nsp16 alone (Kd = 20.0 ± 2.7 μM) (fig. S1D and table S1). We also determined the binding affinities for the methyl donor SAM and the product S-adenosylhomocysteine (SAH) for nsp16 and the nsp16-nsp10 heterodimer (fig. S1, A and B, and table S1). Neither SAM nor SAH bound to nsp16 alone, but both SAM and SAH bound to the nsp16-nsp10 heterodimer with Kd values of
and Mg$^{2+}$ (Fig. 1, C and D). Crystal #2 (PDB 7L6R) was soaked with RNA and the presence of Mg$^{2+}$ or Mn$^{2+}$ stimulate the rates of catalysis. nsp10 greatly enhances the binding affinity. However, a capped short RNA that the Cap-0 analog is capable of binding to nsp16 alone and that nsp10 greatly enhances the binding affinity. However, a capped short RNA and the presence of Mg$^{2+}$ or Mn$^{2+}$ stimulate the rates of catalysis.

Crystal structures of the nsp16-nsp10 heterodimer with capped RNA and metal ions

To gain insight into how metal ions stimulate catalysis, we took a structural biology approach. Crystals of nsp16-nsp10 in complex with SAM from different crystallization conditions were soaked with the custom-synthesized m$^7$GpppAUUAAA substrate in the presence of Mg$^{2+}$ or Mn$^{2+}$. Multiple datasets were collected, and, ultimately, the three with the highest resolution and the best data statistics were selected for further analysis (table S2). Crystal #1 [Protein Data Bank (PDB) 7JYY] was soaked with substrates in the presence of 5 mM MgCl$_2$ for 1.5 hours. In this crystal, we observed Cap-0-RNA, SAM, and Mg$^{2+}$ (Fig. 1, C and D). Crystal #2 (PDB 7L6R) was soaked with substrates in the presence of 20 mM MnCl$_2$ for 6 hours. In this crystal, we observed Mn$^{2+}$ and products of the reaction, Cap-1-RNA and SAH, indicating that the methyl transfer occurred in the crystal (Fig. 1E). Crystal #3 (PDB 7L6T) was soaked with substrates in the presence of 500 mM MgCl$_2$ for 6 hours. In this crystal, we observed two Mg$^{2+}$ ions and the products Cap-1-RNA and SAH. The first Mg$^{2+}$ occupied the same metal binding site as in crystals #1 and #2, and the second Mg$^{2+}$ directly interacted with phosphate groups of the capped RNA (fig. S2). The capped RNAs in crystals #1 and #3 included the m$^7$G cap, the first three ribonucleotides, and the phosphate group of A$_4$. Crystal #2 contained the m$^7$G cap, the first four ribonucleotides, and the phosphate group of A$_5$.

**Substrates are properly aligned when the capped short RNA is present**

The superposition of the previously reported structure of SARS-CoV-2 nsp16-nsp10 with a Cap-0 analog [PDB 6WRZ (25)] and nsp16-nsp10 with Cap-0-RNA (PDB 7JYY, this study) revealed that they are very similar with a root mean square deviation of 0.33 Å (Fig. 2A). We previously showed that the cap-binding site, also called the high-affinity binding site (HBS), is bordered by flexible loops that adopt an open conformation upon Cap-0 analog binding (25). Interactions of the nsp16 residues with the Cap-0 analog and Cap-0-RNA are similar in both structures. The m$^7$GpppA in the HBS is stabilized by stacking of the m$^7$G and A$_1$ bases with Tyr$_{6828}$ and Tyr$_{6930}$ residues, respectively (Fig. 2A). The O2’ of the A$_1$ ribose interacts with the conserved nsp16 catalytic residues (19, 20) as well as with a conserved water molecule we previously identified (25). The interactions between Asn$_{6841}$, SAM, and O2’ from the A$_1$ are also consistent between these structures (Fig. 2B). Of particular note was the space between the Tyr$_{6930}$ and Asp$_{6873}$ side chains, which is occupied by the A$_1$ base in the Cap-0 analog structure. In the Cap-0 RNA structure, this space accommodates the stacked bases of A$_1$ and U$_2$, with the U$_2$ base force repositioning of the A$_1$ ribonucleotide (Fig. 2A). Superposition of the Cap-0 and Cap-0-RNA structures revealed that this repositioning involves both a 0.6-Å shift of the A$_1$ base toward the side chain of Tyr$_{6930}$ without notable changes in the positions of Asn$_{6841}$ and Tyr$_{6930}$, as well as a 0.4-Å decrease in the distance between O2’ of A$_1$ and the SAM methyl group, which occurs without apparent changes in the position of catalytic residues (Fig. 2B). The movement and alignment of the A$_1$ O2’ atom toward the methyl group of SAM may explain why the additional ribonucleotides increase the efficiency of the methyl transfer reaction (33–35).
Although the superposition of Cap-0 and capped RNA structures revealed differences in the position of the first adenosine, no deviations were observed between Cap-0-RNA and Cap-1-RNA conformations (Fig. 2, A and B), indicating that the A₁ base is not repositioned after the methyl transfer. The structures are essentially identical, with the only difference being the methyl group, which moves from SAM to the A₁ ribose hydroxyl group during methyl transfer (Fig. 2, C and D).

### Interaction of capped RNA ribonucleotides with residues of the low-affinity binding site

The structures in complex with capped RNAs also revealed the importance of the low-affinity binding site (LBS) residues for the conformation of the mRNA in the catalytic site. The best resolved and most complete electron density for capped RNA was observed in crystal #2. The position of U₃ in the active site is "locked" by multiple interactions (Fig. 3A). The phosphate group of U₂ interacts with water molecules from the hydration sphere of the metal ion and the side-chain nitrogen of Lys₆₈₇₄; the O₂ of the U₂ base interacts with the main-chain oxygen of Asp₆₈₇₃, and the O₂ of the U₂ ribose makes direct interactions with one of the oxygens of the side chain of Asp₆₈₇₃ (Fig. 3B). The phosphate group of U₃ interacts directly with the side-chain nitrogen of Lys₆₈₇₄ and forms water-mediated interactions with residues Asp₆₈₇₃, Lys₆₈₇₄, Met₆₈₄₀, and Asn₆₈₄₁. The base of U₃ interacts directly with the main-chain oxygen and nitrogen atoms of Ala₆₈₃₂ and forms a water bridge interaction with the nitrogen of the main chain of Leu₆₈₄₄. The whole nucleotide A₄ and phosphate group of A₅, the last ordered part of the capped RNA in the crystal #2 structure, are solvent exposed and connected to the protein by a hydrogen bond interaction between O4’ of the A₄ and the side-chain oxygen of Ser₆₈₃₁ (Fig. 3B). The stacking interactions between bases of U₃ and A₄ define the position of the A₅ nucleotide. It is unknown if the conformation of A₅ reflects the natural interaction of nucleotides or if a longer mRNA would form different contacts with the nsp16-nsp10 heterodimer. However, the position of m₇G and first three nucleotides of the capped RNA closely match in all three structures and likely represent the accurate binding mode for this part of the capped RNA.

### Metal ions stabilize the capped RNA in the nsp16 active site

The primary metal binding site is located near the HBS (Fig. 1C), with either Mg²⁺ or Mn²⁺ occupying the same site with similar interactions (Fig. 3, B and C). The metal ions make both direct and water-mediated interactions with side chains of nsp16 residues and the backbone of the capped RNA. The best electron density maps were observed for crystal #3 with two magnesium ions, both of which have near-ideal octahedral geometry (Fig. 3C). The Mg²⁺ in the primary metal binding site is coordinated in part by interactions with phosphate groups of the triphosphate bridge linking the cap to A₁, the phosphate group of U₂, and the ribose of U₃. The second Mg²⁺ directly interacts with the U₁ and A₄ phosphate group oxygens and through water molecules with the side-chain oxygens of the Asp₆₈₇₃ and the side-chain nitrogen of Lys₆₈₇₄ (Fig. 3C). Thus, the metal ion that occupies the primary metal binding site of nsp16 properly aligns capped RNA with SAM for an efficient methyl transfer reaction. The role of metal ions in facilitating orbital alignments for efficient catalysis has been demonstrated structurally for hydride transfer reactions (34). Structural evidence for the requisite orbital alignment of substrates in RNA 2'-O-MTases (33) and ribozymes (35) has also been observed.

**Asp₆₈₇₃ and Lys₆₈₇₄ alter the backbone conformation of the capped RNA in coronaviruses**

Comparison of the RNA binding site and the capped RNA conformation from the SARS-CoV-2 nsp16-nsp10 structure (PDB 7L6R) and the crystal structures of 2'-O-MTases with capped RNA for DENV NS5 [PDB 5D7O (21)], vaccinia virus (VACV) VP39 [PDB 1AV6 (36)], and human cap MTase hCMTr1 [PDB 4N48 (37)] revealed that the conformation of the m₇G and the location of the cap-binding pockets relative to the active sites are markedly different (fig. S3, A to D). In the nsp16-nsp10 and VP39 structures, the N₇-methyl groups are nestled in the HBS pocket (fig. S3, A and C). In contrast, in NS5 and hCMTr1,
for which methylation at the $N^7$ position of the $G_0$ is not required for the 2′-O-MTase activity (37), the $N$'-methyl groups are pointed toward the solvent (fig. S3, B and D). Although the m′G positions do not overlap, nucleotides $N_1$ and $N_2$ in all these structures are closely matched, which is consistent with the conserved mechanism of action and the structure of the catalytic site (Fig. 4A). In all but the nsp16-nsp10 structure, the $N_2$ nucleotide is sandwiched between the $N_1$ and $N_3$ by base stacking interactions, and these three nucleotides have limited interactions with the 2′-O-MTase residues of RNA binding groove. In the nsp16-nsp10 structure, all functional groups of the nucleotides $N_2$ and $N_3$ are involved in an integrated and complex network of hydrogen bond interactions with residues of the LBS. The $U_2$ base and the ribose directly interact with main-chain and side-chain atoms of Asp$^{6873}$. The Asp$^{6873}$ side chain occupies the space that is filled by $N_3$ in all other structures and forces the $U_3$ nucleotide to move to the opposite side of the RNA binding groove. The base of $U_3$ is involved in direct and water-mediated interactions with Ala$^{6832}$ and Leu$^{6834}$, while the phosphate group interacts with Lys$^{6874}$ (Fig. 3, B and C).

The superposition of SARS-CoV-2 nsp16-nsp10 (PDB 7JYY) with MERS-CoV [PDB SYNM (38)] revealed almost identical conformations of the proteins and the promotory Asp$^{6873}$ and Lys$^{6874}$ residues, as well as the position of the Cap moiety (Fig. 4B). Alignment of the amino acid sequences of nsp16 from representative coronaviruses (Fig. 4C and fig. S4, A and B) showed that Asp$^{6873}$ is conserved across the coronaviruses, except for feline coronavirus (F-CoV). Structural alignment of nsp16 with NS5, VP39, and hCMTr1 revealed that two important residues, Asp$^{6873}$ and Lys$^{6874}$, are located in a four-residue loop that connects $\beta 1$ and $\alpha A$ (Fig. 4D). This four-residue insert is uniquely present in all coronavirus nsp16 MTases and is absent from other MTases (Fig. 4E).

To test the effect of this insert on nsp16 MTase activity, we prepared nsp16 proteins with D6873A or D6873G amino acid substitutions or with a deletion of Asp$^{6873}$ and Lys$^{6874}$ within the loop [Δ(D$^{6873}$-K$^{6874}$)]. Overall, Mn$^{2+}$ stimulated the methyl transferase activity of nsp16 better than did Mg$^{2+}$ for both the wild-type and mutant nsp16 proteins (Fig. 5, A and B). At a 600 nM protein concentration in the presence of Mn$^{2+}$, we observed 10% of the maximum activity for all proteins, whereas only 2% activity was observed in the presence of Mg$^{2+}$. A similar pattern was detected when we measured values for maximum activity, which was reached at an enzyme concentration of 500 nM in the presence of Mn$^{2+}$ but required 1 μM enzyme in the presence of Mg$^{2+}$ (Fig. 5, A and B). The EC$_{50}$ (Half maximal effective concentration) values for activity of the D6873A and D6873G mutants in the presence of Mn$^{2+}$ were shifted higher than values obtained for the wild-type enzyme (Fig. 5C). These shifts in EC$_{50}$ values were more evident in the presence of Mg$^{2+}$. The Δ(D$^{6873}$-K$^{6874}$) mutant showed much lower...
activity compared to the D6873A and D6873G mutants and the wild-type enzyme. At enzyme concentrations as high as 2 μM, the activity of Δ(D6873-K6874) reached only 45 and 65% of that of the wild-type max activity in the presence of Mn2+ or Mg2+, respectively. The results of activity assays support our structural finding regarding the importance of the four-residue loop that is unique for coronavirus nsp16 MTase as well as a preference for stimulation of the methyl transfer reaction by Mn2+.

**DISCUSSION**

In eukaryotes, 5′ capping and subsequent 2′-O-methylation events are essential for RNA metabolism and protection from cell innate immunity responses (39). Coronaviruses and other viruses have evolved entire RNA capping and methylation machineries to modify their own RNAs, thereby increasing the translation of viral proteins and promoting the evasion of host immune responses (7, 40, 41). For this reason, the viral 2′-O-MTases are suitable targets for drug development. Thus far, all proposed modeling of the Cap-RNA orientation in the LBS of nsp16-nsp10 (19, 24, 26, 32) have relied on the crystal structures of DENV NS5, VACV VP39, and hCMTr1, indicating a knowledge gap for drug design.

The metal ion requirement for DENV, SARS-CoV, and MERS-CoV MTase activity is well known (11, 19, 20). However, the binding site of the metal ion and capped RNA has been shown only for the NS5 structure (21). In the NS5 structure, the Mg2+ ion directly interacts with all three phosphate groups that link the m7G with the A1 and thus neutralizes the negative charge of the triphosphate group. Three additional interactions through its hydration sphere with the bases of the capped RNA help maintain stacking interactions and stabilize the single-stranded RNA helix in the A-form. The described Mg2+ is not involved in any direct hydrogen bond interactions with the residues of the 2′-O-MTase domain. In all SARS-CoV-2 nsp16 structures reported here, the metal ions that occupy the primary metal binding site directly interact with Asn6996. This direct interaction “pins” the metal ion to the negatively charged “nest.” By neutralizing the negative charge, metal ions stabilize the unique conformation of the capped RNA and by direct interaction with Asn6996 help to hold the RNA in the RNA binding site for methyl transfer.

Metal ions tested in the activity assays had different effects on the methyl transfer efficiency of SARS-CoV-2 nsp16. Na+ has a lower charge and Ca2+ has longer metal-ligand distances than Mg2+ or Mn2+, which might yield a more extended capped RNA that is weakly attached to the binding site. We suspect that this is why Na+ and Ca2+ poorly stimulated the methyl transfer reaction in our assays. Mg2+ had been considered as the primary catalytic cofactor and is reported as an important additive for nsp16 activity assays (11, 19, 20, 22). The results of our biochemical assays, by contrast, showed that Mn2+ had the highest stimulation effect on the methyl transfer reaction, and maximum activity was reached at lower concentrations of enzyme. It was also reported that Mg2+ ions in micro- to millimolar concentrations can promote the cleavage of nucleic acids (42), suggesting that Mn2+ is preferable compared to Mg2+. In addition, Mn2+ has a higher reported concentration than Mg2+ in the endoplasmic reticulum and Golgi, where the SARS-CoV-2 replication vesicles...
are formed (43, 44). Together, these findings suggest that Mn$^{2+}$ is the natural cofactor for the methyl transfer reaction in SARS-CoV-2 and other coronaviruses. This information could be very helpful in the design of specific inhibitors to target the metal binding site and impair the activity of the complex.

Comparison of the RNA binding sites and the capped RNA conformations from the prior MTase structures of NS5, VP39, and HCMTTr1 with those of nsp16-nsp10 complexes revealed major differences between them. These four structures can be grouped into two classes: the $N'$ of the m$^7$G buried in the binding pocket, as observed in nsp16 and VP39, and $N'$ exposed to the solvent, as observed in NS5 and hCMTr1. Despite these differences, the superposition of these structures revealed a close match for the positions and the conformations of the first and the second nucleotides of capped RNA. This observation is in line with the conserved mechanism of the $2'-O$-MTase reaction. The major differences in the conformations of the RNA are located in the extended part of the capped RNA where it interacts with the residues of the LBS. In NS5, VP39, and hCMTr1, the N1-N2 nucleotides of the capped RNA are stabilized by stacking interactions. In contrast, Asp$^{6873}$ in nsp16 alters the position of the N3 and prevents the stacking of N2 to N1-N2, and Lys$^{6874}$ stabilizes the position of the U1 phosphate. All functional groups of the nucleotides N2 and N3 are involved in an integrated and complex network of hydrogen bond interactions with residues of the LBS. These distinct differences were found to affect catalysis with mutation of Asp$^{6873}$, leading to a reduced EC$_{50}$ and deletion of Asp$^{6873}$ and Lys$^{6874}$ from the loop shifting the EC$_{50}$ to the micromolar range.

A key advance of this work was the use of a custom-synthesized substrate RNA with ribonucleotides matched specifically to the sequences of the SARS-CoV-2 mRNA 5'-ends compared to earlier studies that used poly(C) RNA (11, 19). Use of this substrate, however, introduced a limitation of the studies due to precious amounts of the compound, which prevented maximizing the substrate concentration for kinetic and binding assays. However, we were able to determine the crystal structure of nsp16-nsp10 in complex with Cap-0-RNA and the product of the reaction, Cap-1-RNA. Cap-0-RNA was used to conduct biochemical assays at a fixed concentration of substrates to show the preference of the enzyme for Mn$^{2+}$ over Mg$^{2+}$, a requirement for the extended RNA, and the importance of the unique promontory loop in nsp16 for catalysis. These studies in total indicate that redirection of the RNA, which is also coordinated by metals, leads to proper alignment of substrates for efficient methyl transfer to the 2'-OH of the first adenosine, and that the presence of a unique four-residue insert (D$^{6873}$K$^{6874}$G$^{6875}$V$^{6876}$), which is conserved in all coronaviruses is critical for catalysis. The absence of this insert in mammalian MTases makes this region a promising site for the design of selective RNA-like coronavirus-specific inhibitors.

**MATERIALS AND METHODS**

**Protein expression, purification, and crystallization**

Recombinant nsp16 and nsp10 proteins were purified from *Escherichia coli* and crystallized as previously described (25). DNA sequences for expression of mutant nsp16 protein were also generated by synthetic DNA in the same vector (Twist Biosciences) and purified following the prior established protocol (25). To form the nsp16-nsp10 heterodimers, the pure proteins were mixed at a 1:1 molar ratio at ~2 mg/ml in buffer [10 mM tris-HCl (pH 8.3), 500 mM NaCl, 1 mM tris(2-carboxyethyl) phosphine (TCEP), 2 mM MgCl$_2$ and 5% glycerol] and incubated for 1 hour, then dialyzed in crystallization buffer [10 mM tris-HCl (pH 7.5), 150 mM NaCl, 2 mM MgCl$_2$, 1 mM TCEP, and 5% glycerol] for 2 hours. SAM was added to a final concentration of 2 mM. Freshly purified preformed nsp16-nsp10 heterodimer was concentrated to 4.0 to 5.5 mg/ml and immediately set up for crystallization as 2-μl crystallization drops (1 μl protein:1 μl reservoir solution) in 96-well crystallization plates.

**MTase activity assays**

Custom-synthesized Cap-0-RNA (m$^7$GpppAUUAAA) was obtained from Bio-Synthesis Inc. (Lewisville, TX). Cap-0 analog (m$^7$GpppA) was obtained from New England Biolabs (catalog #SI405L). The MTase activity was measured using the MTase-Glo Methyltransferase bioluminescence assay (Promega) (45) according to the manufacturer’s protocol in a suitable solution for SARS-CoV-2 MTase: 20 mM tris-HCl (pH 8.0), 1 mM EDTA, 100 nM nsp16-nsp10 heterodimer, 1 μM excess of nsp10, 5 μM SAM, 300 nM Cap-0-RNA or Cap-0 analog, and 2 mM MgCl$_2$, MnCl$_2$, CaCl$_2$, or 3 mM NaCl as indicated. The reactions were incubated for 1 hour at 37°C and stopped with 0.5% trifluoroacetic acid. The detection solution from the kit was then added, and the mixture was further incubated for 30 min at room temperature, followed by the addition of the developing solution. Luminescence was measured using a TECAN Safire2 microplate reader in arbitrary units and normalized assigning 100% to the activity in the presence of Mg$^{2+}$ and Cap-0-RNA. The average and the SD of three measurements in two independent experiments using different protein purifications ($n = 6$) were plotted as a histogram using GraphPad Prism v9.

For determination of EC$_{50}$, the assay was modified for a 384-well plate format at the Northwestern University High Throughput Analysis Laboratory. Each nsp16-nsp10 heterodimer complex was serially diluted in solution as detailed above using the Mosquito robot. Reactions were initiated by addition of the substrates (5 μM SAM and 1 μM Cap-0-RNA) using the Mantis liquid handler. After 1 hour at 37°C, the reactions were coupled with the detection and developing solutions using Mantis liquid handler and incubated as described above. Luminescence was measured in a TECAN Infinite M1000. The blank was defined as containing all reagents except Cap-0-RNA, and this value was subtracted for each corresponding point. The values were then reported as the percentage of MTase activity compared to the wild type under the same conditions. Assays were conducted in triplicate, and the experiment was independently repeated ($n = 6$). EC$_{50}$ and the standard error were calculated using four-parameter logistic curve in GraphPad Prism v9 at 95% confidence interval.

**Isothermal titration calorirometry**

Binding affinity was determined using a MicroCal PEAQ-ITC system (Malvern, Worcestershire, UK) at 25°C. The sample cell volume was 200 μl, and the total syringe volume was 40 μl. For each titration, the first injection was performed using 0.4 μl, which was then followed by 18 additional injections at 2 μl per injection. The first injection was considered a void and was automatically removed from data analysis. Each injection was spaced by 120 s after a 60-s initial delay. SARS-CoV-2 samples of nsp10, nsp16, and nsp16-nsp10 were individually loaded into the sample cell and then titrated with either SAH, SAM, m$^7$GpppA Cap analog or the m$^7$GpppG Cap analog.
parameters for \( N \), model and nonlinear curve fitting. Each experiment was performed with MicroCal PEAQ-ITC Analysis Software using a single-site binding model and nonlinear curve fitting. Each experiment was performed in triplicate, and the resulting values and standard error in the fitted parameters for \( N, K_d, \Delta H, \Delta S, \) and \( AG \) were obtained and are summarized in table S2.

Crystal soaking experiments

Diffraction quality crystals grown from different conditions were transferred into 10-\( \mu \)l drops containing 5 mM SAM, 0.2 mM Cap-0-RNA, and into 10-\( \mu \)l drops with 5 mM SAM, 0.2 M Cap-0-RNA, and 5 mM MgCl\(_2\) or 20 mM MnCl\(_2\) in respective reservoir solutions and soaked for different time periods from 5 min to 17 hours. Crystals were cryoprotected with 4 mM sodium formate or 25% sucrose in the respective reservoir solution and flash frozen in liquid nitrogen for data collection.

Crystal #1 was grown from 0.1 M sodium citrate (pH 5.6), 1.0 M ammonium dihydrogen phosphate, soaked for 1.5 hours with 0.2 mM Cap-0-RNA, 5 mM SAM, 5 mM MgCl\(_2\), and cryoprotected with 4 M sodium formate. Crystal #2 was grown from 0.1 M citric acid (pH 5.0), 0.8 M ammonium sulfate, and was soaked for 6 hours with 0.2 mM Cap-0-RNA, 5 mM SAM, and 20 mM manganese chloride. Crystal #3 was grown from 0.1 M Hepes (pH 7.5), 0.5 M magnesium formate and was soaked for 6 hours with 0.2 mM Cap-0-RNA and 5 mM SAM. Crystals #2 and #3 were cryoprotected with 25% sucrose in the respective reservoir solutions.

Data collection, processing, structure solution, and refinement

More than 30 crystals were screened, and 32 datasets were collected at the beam lines 21ID-D and 21ID-F of the Life Sciences–Collaborative Access Team (LS-CAT) at the Advanced Photon Source, Argonne National Laboratory. Images were indexed, integrated, and scaled using HKL-3000 (46).

Data quality and structure refinement statistics are shown in table S2. All structures were determined by Molecular Replacement with Phaser (47) from the CCP4 Suite (48) using the crystal structure of the nsp16-nsp10 heterodimer from SARS-CoV-2 as a search model (PDB 6W4H). The initial solutions went through several rounds of refinement in REFMAC v. 5.8.0266 (49) and manual model corrections using Coot (7). The water molecules were generated using ARP/WARP (50) followed by additional rounds of refinement in REFMAC. All structures were carefully examined, and three datasets were selected for further structural studies. For all structures, the Cap-0-RNA, SAM, and Mg\(^{2+}\) were fit into electron density maps and further refined. Inspection of anomalous and Fourier difference electron density maps revealed that, for crystal #1, the nsp16-nsp10 heterodimer formed the complex with the Cap-0-RNA, SAM, and Mg\(^{2+}\); for crystal #2, the complex was formed with the Cap-1-RNA, SAH, and Mn\(^{2+}\); and for crystal #3, the complex was formed with the Cap-1-RNA, SAH, and two Mg\(^{2+}\). In crystals #1 and #3, no additional electron density was detected beyond phosphate group of A\(_1\). In crystal #2, the presence of well-defined electron density near the phosphate group of A\(_1\) allowed the RNA model to be unambiguously extended by adding sugar and base for A\(_1\) and phosphate group of A\(_2\). All structures were further refined with the Translation-Libration-Screw (TLS) group corrections, which were created by the TLS Motion Determination (TLSMD) server (51). The quality control of the models during refinement and for the final validation of the structures was done using MolProbity (52) (http://molprobity.biochem.duke.edu/). The capped RNA has a 5′-5′ triphosphate linkage between m\(^7\)G and A\(_1\) with noncanonical backbone geometry, which resulted in low RNA backbone scores in MolProbity reports. All structures were deposited to Validated SARS-CoV-2-related structural models of potential drug targets (https://covid19.bioreproducibility.org/) and to the Protein Data Bank (www.rcsb.org/) with the assigned PDB 7IYY (crystal #1), 7L6R (crystal #2), and 7L6T (crystal #3). Flexible parts of the structure, alternative side-chain conformations, and partial water molecules were fit into electron density maps at a lower \( \sigma \) level, which resulted in overall high real-space R-value Z-score (RSRZ) scores for all structures. All figures with models of the structures were created in PyMOL open source version V 2.1 (53); the diagram showing protein and capped RNA interactions was created in LigPlot+ (54).

Structural, sequence alignment, and phylogenetic analysis

The PDB coordinates of SARS-CoV-2 nsp16 and nsp10 were analyzed using the FATCAT (55), POSA (56), and DALI (57) servers to perform structural alignments with MERS-CoV, NS5, hCMTr1, and VP39 cap RNA MTases. Generated PDB files were downloaded from the servers and modeled in PyMOL open source version v2.1. The protein sequence of 2′-O-MTases was obtained from the National Center for Biotechnology Information database: F-CoV (AGT52079), murine hepatitis virus (MHV; YP_009915686.1), human coronavirus (HKU1; YP_460023.1), human coronavirus 229E (H-CoV-229E; AGT21344.1:6464-6763), infectious bronchitis virus (IBV; NP_066134.1: 6328-6629), DENV nonstructural protein NS5 (NS5, NP_739590.2), Homo sapiens (hCMTr1, BAA07893.3 KIAA0082), and VACV VP39 (VP39, NC_006998.1). The multiple sequence alignment was performed using Clustal-O (www.ebi.ac.uk/Tools/msa/clustalo/) and merged with the coordinates of the structure with PDB code 7IYY using ESPript 3.x (58). The phylogenetic tree was created using MacVector and processed in iTol (https://itol.embl.de).

SUPPLEMENTARY MATERIALS

stke.sciencemag.org/cgi/content/full/14/689/eabh2071/DC1
Figs. S1 to S4
Tables S1 and S2
References (59–60)

View/request a protocol for this paper from Bio-protocol.

REFERENCES AND NOTES

Coronavirus nsp10/nsp16 methyltransferase can be a protein nsp10 can suppress the intermediate state in SARS-CoV-2 mRNA capping and methylation.  

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**References**


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Mn$^{2+}$ coordinates Cap-0-RNA to align substrates for efficient 2′-O-methyl transfer by SARS-CoV-2 nsp16

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Uniquely coronavirus
Virally encoded 2′-O-methyltransferases catalyze the last step in the capping of viral RNAs, which protects the RNAs from degradation and prevents them from triggering host defenses. Minasov et al. report structures of the SARS-CoV-2 methyltransferase, a heterodimeric complex of the enzyme nsp16 and its coactivator nsp10, in complex with a short, capped RNA (instead of the RNA cap analogs used to generate previous structures), the methyl donor SAM, and divalent metal cations. The metal ions and a four-residue insert of nsp16 were important for precisely aligning the RNA substrate in the active site for efficient catalysis. This insert is present in coronavirus but not in mammalian methyltransferases, suggesting this site as a potential target for the design of coronavirus-specific methyltransferase inhibitors.