2′-C-methylated nucleotides terminate virus RNA synthesis by preventing active site closure of the viral RNA-dependent RNA polymerase

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The 2′-C-methyl ribonucleosides are nucleoside analogs representing an important class of antiviral agents, especially against positive-strand RNA viruses. Their value is highlighted by the highly successful anti-hepatitis C drug sofosbuvir. When appropriately phosphorylated, these nucleotides are successfully incorporated into RNA by the virally encoded RNA-dependent RNA polymerase (RdRp). This activity prevents further RNA extension, but the mechanism is poorly characterized. Previously, we had identified NMR signatures characteristic of formation of RdRp–RNA binary and RdRp–RNA–NTP ternary complexes for the poliovirus RdRp, including an open-to-closed conformational change necessary to prepare the active site for catalysis of phosphoryl transfer. Here we used these observations as a framework for interpreting the effects of 2′-C-methyl adenosine analogs on RNA chain extension in solution-state NMR spectroscopy experiments, enabling us to gain additional mechanistic insights into 2′-C-methyl ribonucleoside-mediated RNA chain termination. Contrary to what has been proposed previously, poliovirus RdRp that was bound to RNA with an incorporated 2′-C-methyl nucleotide could still bind to the next incoming NTP. Our results also indicated that incorporation of the 2′-C-methyl nucleotide does not disrupt RdRp–RNA interactions and does not prevent translocation. Instead, incorporation of the 2′-C-methyl nucleotide blocked closure of the RdRp active site upon binding of the next correct incoming NTP, which prevented further nucleotide addition. We propose that other nucleotide analogs that act as nonobligate chain terminators may operate through a similar mechanism.

RNA viruses represent some of the greatest threats to global health systems, including positive-strand RNA viruses like those belonging to the Flaviviridae (1) and Picornaviridae (2, 3) families. Although vaccines have been or are currently being developed against many of these viruses (4, 5), there has also been a recognized need for small-molecule therapeutic agents to complement or even overtake these approaches (6, 7). One important drug target is the viral RNA-dependent RNA polymerase (RdRp)2 (8, 9), responsible for virus RNA replication. Replication inhibitors that target the conserved core of the RdRp may have broad antiviral activity. For example, 2′-C-methyl nucleotide analogs have wide antiviral activity against many flaviviruses, picornaviruses, and other positive-strand RNA viruses (10–12), although they may have limited utility against negative-strand RNA or DNA viruses (13). This class includes the highly successful anti-hepatitis C drug sofosbuvir (14). The RdRp incorporates the triphosphorylated 2′-C-methyl nucleotide analogs into RNA, but further RNA extension is prevented, despite these molecules containing a 3′-hydroxyl necessary for further nucleotide incorporation (10, 15, 16); such molecules are known as nonobligate chain terminators. The antiviral activity of these compounds is likely due to RNA chain termination (10, 13, 16), but the molecular mechanisms behind RNA chain termination are poorly understood. A better appreciation of these molecular mechanisms may provide insights into further development of this important class of antiviral agents.

The RdRp three-dimensional structure has been described as a cupped right hand with fingers, thumb, and palm subdomains, similar to the overall architecture of other single-subunit nucleic acid polymerases (17). Viral RdRps are unique compared with other nucleic acid polymerases in that there are extensive interactions between the fingers and thumb subdomains, providing completely encircled active sites (18) (Fig. 1). The enclosure of the RdRp active site means that mechanisms of nucleotide selection, addition, and translocation may differ from other nucleic acid polymerases. There are available X-ray crystal structures for a number of RdRps, including those for flaviviruses and picornaviruses (18). Unfortunately, the only X-ray crystal structures with RdRp (in this case, from hepatitis C virus (HCV)) in complex with 2′-C-methyl nucleotides represent a step before nucleotide incorporation, with the triphosphorylated 2′-C-methyl nucleotide acting as the next incoming NTP (19). Although these structures have revealed molecular determinants important for nucleotide analog recognition, they provide little insight into the mechanism of RNA termination. A series of X-ray crystal structures of poliovirus (PV) and related RdRps provides insights into the steps of nucleotide incorporation and subsequent translocation (20–22), including

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This article contains Figs. S1 and S2 and Table S1.

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The abbreviations used are: RdRp, RNA-dependent RNA polymerase; HCV, hepatitis C virus; PV, poliovirus; 2′-C-A, 2′-C-methyladenosine; HSQC, heteronuclear single quantum coherence.

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insights into the functional roles of the seven conserved structural motifs in RdRs (i.e., motifs A–G). After base-pairing with the template nucleotide, the 2′- and 3′-hydroxyls of the incoming NTP make key hydrogen bonding interactions with Ser-288 and Asn-297 in motif B (PV numbering), triggering structural rearrangements to “close” the active site and prepare for the phosphoryl transfer reaction (21, 22). Critically, structural motif A is realigned to form a complete three-stranded β-sheet with motif C, which causes motif A residue Asp-233 to swing toward the RNA, allowing it to coordinate the magnesium ions necessary for catalysis (21). The triphosphate of the incoming NTP is also realigned to promote inline nucleophilic attack from the primer 3′-hydroxyl, aided in part by conserved residues in motifs D and F (21–23). The phosphoryl transfer reaction then takes place, according to the two metal ion mechanism shared by other nucleic acid polymerases (24), with Asp-233 and motif C residue Asp-328 coordinating the metals (21). We have also proposed that there is a repositioning of motif D residue Lys-359 so that it can act as a general acid to protonate the pyrophosphate leaving the group (25–27). Following nucleotide addition, the active site reopens to release the pyrophosphate, but this conformational change is not tightly coupled to translocation as it is in other nucleic acid polymerases (22, 28). It has been proposed that additional structural changes in motif B then mechanically aid translocation (29, 30) to eventually reset the protein to accept the next incoming NTP.

The 2′-C-methyl ribonucleotides may act at any of these steps to terminate RNA synthesis. Initial modeling studies have proposed that the next incoming NTP is prevented from binding because of a steric clash with the 2′-C-methyl group (13). Excision of the incorporated 2′-C-methyl nucleotide through pyrophosphorolysis (i.e., the reverse of the nucleotide addition reaction) was not sensitive to the next NTP, which, the authors suggested, was also evidence that NTP binding was impeded (31). Incorporation of the 2′-C-methyl nucleotide might also disrupt interactions between the RdRp and RNA to promote RNA release, diminish the ability of the RdRp to form the catalytically competent closed conformation, and/or inhibit translocation after nucleotide analog incorporation.

In the absence of X-ray crystal structure data, we used solution-state NMR studies of PV RdRp, along with steady-state kinetics, to provide insight into the mechanism of RNA chain termination by 2′-C-methyl ribonucleotides. These studies suggest that incorporation of 2′-C-methyl nucleotide prevents subsequent active-site closure, a prerequisite step for incorporation of the next incoming nucleotide.

**Results**

**Incorporation of 2′-C-Me-AMP by PV RdRp terminates RNA synthesis and inhibits virus replication**

We first tested the ability of PV RdRp to incorporate 2′-C-methyl nucleotide. Here we used 2′-C-methyladenosine (2′-C-Me-A) as a representative member of the 2′-C-methyl nucleotide analogs (Fig. 2), and the ssAU RNA, which contains a 6-bp duplex flanked by four nucleotide 5′ overhangs (i.e., 5′-GCAU-
previously used [methyl-\(^{13}\)C]Met groups in PV RdRp to gain insight into structural and dynamic changes important for nucleotide selection and catalysis (27, 35). The [methyl-\(^{13}\)C]Met probes are conveniently located in or near regions that have been shown by X-ray crystallography structures and/or molecular dynamics simulations to undergo structural and/or dynamic changes upon binding RNA and/or NTP (20–22, 36, 37) (Fig. 1). In this study, we were especially interested in the Met-6, Met-74, Met-187, Met-225, Met-354, and Met-394 probes, as their associated resonances have been shown to undergo chemical shift and/or peak intensity changes upon binding RNA and/or NTP (27, 35) (Fig. 3). Other Met probes are more solvent-exposed and so do not respond to ligand binding and/or associated structural changes (35).

Met-6, Met-74, and Met-187 are located in the fingers subdomain and likely respond to structural rearrangements in motifs A, B, and C. Met-6 is part of the N-terminal \(\beta\)-strand that makes hydrogen bond interactions with palm structural motifs A and B. Met-6 is also part of the three-strand \(\beta\)-sheet that makes contact with residues in structural motif F (Fig. 4). Motif F is important for RNA template and incoming NTP binding (20, 21). Arg-174 in motif F is likely responsible for properly aligning the \(\alpha\)-phosphate with the primer 3' hydroxyl (38); the R174K-substituted variant provides the highest nucleotide selection incorporation fidelity for any PV variant characterized so far (32), and Coxsackievirus encoding the R174K substitution is not viable (39). His-273 is also part of this interaction network, and it has been shown that the H273R substitution results in a lower nucleotide selection fidelity; PV encoding the H273R substitution has reduced virulence in a mouse model (36, 40). Met-74 makes van der Waals contact with the motif B \(\alpha\)-helix (Fig. 4) and with residues on the e1 \(\alpha\)-2 loop that also interact with motifs A and B. Met-187 makes contact with the motif B loop important for recognition of the ribose of the incoming NTP (Fig. 4), and residues in this loop may be important for inducing translocation along the RNA (29).

Met-225 and Met-354 are both located in the palm subdomain. Met-225 makes contact with residues in motifs A and C (Fig. 4) and likely responds to structural changes in these regions, including realignment of the \(\beta\)-strand in motif A as part of active-site closure (21) (Fig. 1). Met-354 is in motif D (Fig. 4), and we have proposed that this probe is responsive to structural changes that reposition the general acid Lys-359 for catalysis (27). This probe might also be responsive to accompanying structural changes in motifs A and B. Met-394 is on the thumb subdomain, near the nascent RNA channel (Fig. 4), and likely responds on structural changes important for binding and positioning the RNA primer/template.

Our previous NMR studies of RdRp–RNA binary and RdRp–RNA–NTP ternary complexes provide a framework for comparing this study with the 2'-C-Me-A analogs. The \(^{1}H\) and \(^{13}C\) chemical shifts for all complexes analyzed in this manuscript can be found in Table S1. Upon addition of ssAU RNA to ligand-free PV RdRp, there were chemical shift changes to the Met-6, Met-187, Met-225, Met-354, and Met-394 resonances and a peak intensity change in the Met-74 resonance (35), suggesting structural and/or dynamics changes in all of the associ-
NMR studies of 2′-C-methylated nucleotide

Figure 3. (Methyl-13C)Met chemical shift perturbations provide insight into structure and dynamic changes in PV RdRp upon binding RNA and nucleotide. A, 1H-13C HSQC of PV RdRp in the absence of RNA and nucleotide. Important Met resonances are labeled. Other Met resonances are not shown. Chemical shifts of RdRp in the presence or absence of RNA or nucleotide were not observed. B–D, 1H-13C HSQC comparisons between PV RdRp in the absence of RNA and nucleotide (black), PV RdRp bound to ssAU(3′dA) RNA (blue), PV RdRp bound to ssAU(3′dA) RNA and UTP (red), and PV RdRp bound to ssAU(3′dA) RNA and CTP (green). The resonances for Met-187 are indicated in the insets. For these experiments, PV RdRp (250 μM) is first incubated with 500 μM duplex RNA (ssAU (Fig. 2)) and 4 mM 3′-dATP so that 3′-dAMP is incorporated but lack of the 3′-hydroxyl prevents further nucleotide addition. Excess 3′-dATP is removed through a desalting column before addition of the second NTP (4–12 mM) to generate the ternary RdRp–RNA–NTP complexes. The D2O-based buffer consisted of 10 mM HEPES (pH 8.0), 200 mM NaCl, 0.02% NaN3, 5 mM MgCl2, and 10 μM ZnCl2. NMR spectra were collected at 293 K using a Bruker Avance III 600 MHz spectrometer. Aspects of these NMR experiments have been reported previously (27, 35, 36).

ated regions (Fig. 3 and Fig. S1). To gain insight into ternary RdRp–RNA–NTP complexes, we first added 3′-dATP, which the RdRp incorporates into the ssAU RNA to produce the ssAU(3′dA) RNA, and then the RdRp–RNA complex was passed through a desalting column to remove excess 3′-dATP. Previous studies indicated minor chemical shift differences for Met-187 and Met-354 between the RdRp–ssAU and RdRp–ssAU(3′dA) binary complexes (35) (Fig. S1). The RdRp–ssAU(3′dA) RNA complex can bind but not incorporate the next incoming NTP. Addition of the next correct NTP to form the RdRp–ssAU(3′dA)–UTP ternary complex resulted in additional chemical shift changes in the Met-6, Met-74, Met-187, Met-225, and Met-354 resonances (Fig. 3 and Fig. S1). Addition of incorrect NTP (2′-dUTP or CTP) resulted in chemical shift changes to the Met-6, Met-187, Met-354, and Met-394 resonances compared with what was observed for the RdRp–RNA(3′dA) binary complex, indicating that the incorrect NTP still binds but does not induce the same conformational changes as the correct NTP (Fig. 3). Notably, the chemical shift patterns were different between the RdRp–ssAU(3′dA)–UTP ternary complex and the RdRp–ssAU(3′dA)–2′-dUTP and RdRp–ssAU(3′dA)–CTP ternary complexes (Figs. 3 and 5). These studies established NMR “fingerprints” for when the enzyme is in an “open” (e.g. RdRp–ssAU(3′dA)–CTP) or a closed (e.g. RdRp–ssAU(3′dA)–UTP) conformation. In the open conformation, it is likely that Asp-233, the triphosphate of the incoming NTP, and/or the general acid Lys-359 are not properly positioned for the nucleotide addition reaction.

The RdRp–RNA ternary complex with 2′-C-methyl-ATP does not achieve the closed conformation

To better understand how 2′-C-methyl-ATP interacts with PV RdRp, binary and ternary complexes were evaluated using the ssUU RNA (5′-GCUUGGGCCC-3′), which was 3′-H-terminated according to previous procedures. There were some differences in the NMR spectra between the RdRp–ssUU(3′dA) and RdRp–ssUU(3′dA) binary complexes (Fig. 5). In particular, there were minor chemical shift differences for the Met-225 and Met-354 resonances. Perhaps more interestingly, there was evidence of conformational exchange in regions associated with Met-187 and Met-394. Met-394 was associated with two resonances, and the Met-187 resonance was at very low intensity, suggestive of conformational exchange on the intermediate NMR timescale. These findings suggested that the RdRp complexes with ssUU(3′dA) were more structurally dynamic, which complicated spectral analysis.

The ternary complex with correct incoming NTP (i.e. RdRp–ssUU(3′dA)–ATP) also exhibited signs of conformational exchange (Fig. 5). In this case, there were (at least) two resonances associated with Met-225, Met-354, and Met-394. The two resonances for Met-354 in the RdRp–ssUU(3′dA)–ATP complex had similar chemical shift positions as those observed...
for Met-354 in the RdRp–ssAU(3’dA)–UTP and RdRp–ssAU(3’dA)–CTP complexes. This result suggested that the RdRp–ssUU(3’dA)–ATP complex fluctuated between the closed and open states, although, in this case, it appeared that the open state was more favored. Although this result was not expected, it should be noted that previous single-nucleotide incorporation studies indicated that AMP incorporation templated against U had the lowest $k_{pol}/K_{d, app}$ value for all “correct” nucleotide incorporations (41). This result may be due to a more limited ability of PV RdRp to fluctuate into the more active closed conformation, consistent with the results observed here. NMR studies with other RNA templates may provide more insight. Nonetheless, the Met-187 resonance for the RdRp–ssUU(3’dA)–ATP complex had a very low peak intensity, again suggestive of conformational exchange, although the chemical shift position was different from that for the RdRp–ssUU(3’dA) complex (Fig. 5). The Met-225 probe did not show evidence of conformational exchange. In contrast, both Met-354 and Met-394 were associated with two (or more) resonances, but neither of the Met-354 resonances corresponded to chemical shift positions characteristic of the closed conformation, as observed for the RdRp–ssAU(3’dA)–UTP and RdRp–ssUU(3’dA)–ATP complexes.

**A mispaired RNA terminus prevents active-site closure**

Because we have established that PV RdRp can successfully incorporate 2’-C-Me-AMP but then cannot extend RNA synthesis (Fig. 2), we wanted to better understand the downstream consequences of 2’-C-Me-AMP incorporation. We had previously analyzed the consequences of GMP misincorporation (templated against U) (27). For the NMR studies, we incubated ssAU RNA with 3’-dGTP and then removed excess 3’-dGTP by passage across a desalting column to generate the RdRp–ssAU(3’dA) binary complex. We did not note substantial differences between the RdRp–ssAU(3’dA) and RdRp–ssAU(3’dG) complexes (Fig. 6), except for the Met-187 resonance. This resonance had a very low peak intensity for the RdRp–ssAU(3’dG) complex, suggesting conformational exchange on the intermediate NMR timescale for the associated region. The small differences between the RdRp–ssAU(3’dA) and RdRp–ssAU(3’dG) complexes suggested that these complexes reached similar conformations, except perhaps in or near motif B.

There were additional chemical shift changes for the Met-187, Met-354, and Met-394 resonances upon addition of the

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**Figure 4. Methionine probes report on ligand binding and conformational changes throughout the RdRp structure.** The template and primer RNA strands are colored red and blue, respectively. **A**, Met-6 is part of the three-stranded β-sheet that makes interactions with residues in motif F (pink). **B**, Met-74 is in the fingers subdomain and makes van der Waals contact with residues on the motif B helix (green). Met-354 is in motif D (blue). Other important residues include the proposed general acid Lys-359 and Asn-297, which makes hydrogen-bonding interactions with the 2’-hydroxyl of the incoming NTP. **C**, Met-187 is near motif B, including Ser-288 and Asn-297, and so likely reports on RNA and NTP binding. **D**, Met-225 is near motifs A and C (red and yellow, respectively) and likely reports on the realignment of motif A to form the three-stranded β-sheet important in active-site closure. **E**, Met-394 is near residues important for RNA binding.
NMR studies of 2’-C-methylated nucleotide

Figure 5. The RdRp–RNA ternary complex with 2’-C-Me-ATP does not achieve the closed conformation. A, experimental design. PV RdRp (250 μM) is first incubated with 500 μM duplex RNA (ssAU or ssUU) and 4 mM 3’-dATP so that 3’-dAMP is incorporated, but lack of the 3’-hydroxyl prevents further nucleotide addition. Excess 3’-dATP is removed through a desalting column before addition of the second NTP (4–12 mM) to generate the ternary RdRp–RNA–NTP complexes. The D2O-based buffer consisted of 10 mM HEPES (pH 8.0), 200 mM NaCl, 0.02% NaN3, 5 mM MgCl2, and 10 μM ZnCl2, B–H, [13C-methyl]Met 1H-13C HSQC NMR spectra of different RdRp–RNA binary and RdRp–RNA–NTP ternary complexes, including the RdRp–ssAU(3’dA) (i.e. ssAU RNA with incorporated 3’-dAMP) (B), RdRp–ssAU(3’dA)–UTP (C), RdRp–ssAU(3’dA)–2’-dUTP (D), RdRp–ssAU(3’dA)–CTP (E), RdRp–ssUU(3’dA)–2’-C-Me-ATP (F), RdRp–ssUU(3’dA)–ATP (G), and RdRp–ssUU(3’dA)–2’-C-Me-ATP (H) complexes. Resonances belonging to the ε1CH3 groups of Met-187, Met-225, Met-354, and Met-394 are highlighted. NMR spectra were collected at 293 K using a Bruker Avance III 600 MHz spectrometer.

next correct nucleotide to form the RdRp–ssAU(3’dG)–UTP ternary complex (Fig. 7 and Fig. S1). The chemical shift positions for the Met-6, Met-74, Met-225, Met-354, and Met-394 resonances were very similar between RdRp–ssAU(3’dA)–CTP (i.e. matched RNA terminus, incorrect incoming NTP) and RdRp–ssAU(3’dG)–UTP (i.e. mismatched RNA terminus, correct incoming NTP), suggesting that the RdRp–ssAU(3’dG)–UTP complex also did not form the closed conformation (Fig. S1). Consistent with this result was the finding that nucleotide incorporation onto RNA with a terminal mispair was slowed to a similar extent as incorrect nucleotide incorporation onto RNA with a properly paired terminus (Fig. 2).

2’-C-Me-AMP incorporation has minor effects on the RdRp–RNA binary complex

We collected NMR spectra for additional RdRp–RNA binary and RdRp–RNA–NTP ternary complexes with mismatched RNA termini by incorporating 2’,3’-ddATP (i.e. ssAU(2’3’ddA)) and 2’-C-methyl-ATP (i.e. ssAU(2’CA) and ssUU(2’CA)) (Figs. 6 and 7 and Fig. S2). The RdRp–ssAU(2’3’ddA), RdRp–ssAU(2’CA), and RdRp–ssUU(2’CA) binary complexes had similar NMR spectra and only minor differences compared with the NMR spectra of the RdRp–ssAU(3’dA) and RdRp–ssUU(3’dA) complexes (Fig. 6). Along with the Met-354 resonance observed for the RdRp–ssAU(3’dA) and RdRp–ssUU(3’dA) complexes, there was another lower-intensity resonance observed for the RdRp–ssAU(2’3’ddA), RdRp–ssAU(2’CA), and RdRp–ssUU(2’CA) complexes, suggesting some conformational exchange in motif D and associated regions. There were also two resonances associated with Met-394, although similar behavior was observed for the RdRp–ssUU(3’dA) complex. Perhaps most interesting, the Met-187 resonance for the RdRp–ssAU(2’CA) complex was more intense than that observed in either the RdRp–ssAU(3’dG) and RdRp–ssUU(2’3’ddA) complexes, although at a slightly different chemical shift position than that observed for the RdRp–ssAU(3’dA) complex. This finding suggests that motif B and surrounding regions are less conformationally dynamic when the RdRp is bound to ssAU(2’CA) compared with when bound to either ssAU(3’dG) or ssUU(2’3’ddA). The incorporated 2’-C-Me-AMP may still make appropriate interactions with the RdRp, including a hydrogen bond between its 2’-hydroxyl and the backbone carbonyl Tyr-326, to limit conformational exchange in this region.

Incorporation of 2’-C-Me-AMP prevents active-site closure

The NMR spectra for the RdRp–RNA–NTP ternary complexes with mismatched RNA termini provided insight into 2’-C-Me-AMP–induced chain termination (Fig. 7). The NMR spectrum for the RdRp–ssAU(2’CA)–UTP complex was very similar to that of the RdRp–ssAU(3’dG)–UTP and RdRp–ssAU(3’dA)–CTP complexes. The RdRp–ssUU(2’CA)–UTP complex also had similar resonances, although analysis of this spectrum was complicated by conformational exchange events observed in the ssUU-bound complexes that appeared to be absent in the ssAU-bound complexes. These results imply that UTP was capable of binding to the RdRp–ssAU(2’CA) and RdRp–ssUU(2’CA) complexes but that the ternary complex...
was not able to fluctuate into the closed conformation necessary for nucleotide incorporation.

The next incoming NTP can still bind after 2'-C-Me-AMP incorporation

Our NMR studies suggested that the next incoming NTP could still bind after 2'-C-Me-AMP incorporation, although the ternary complex is not able to fluctuate into the closed conformation necessary for the phosphoryl transfer reaction. This proposal is in contrast to the previous suggestion that 2'-C-Me-AMP incorporation sterically prevents the next NTP from binding (13). To verify our proposal, we conducted steady-state nucleotide incorporation assays (32) to monitor the effects of the next incoming NTP on RdRp–RNA complex dissociation (Fig. 8). Under steady-state conditions, the rate-determining step for nucleotide incorporation is dissociation of the RdRp–RNA_n+1 complex so that the RdRp can bind to another RNA_n for another nucleotide incorporation reaction (32, 42) (Fig. 8A). Addition of the next incoming NTP may have an effect on the rate of RdRp–RNA complex dissociation (Fig. 8). For example, the steady-state incorporation of 3'-dAMP was reduced in the presence of UTP, likely because the RdRp becomes trapped in a tight closed conformation (i.e. RdRp–ssAU[3’dA]–UTP) that prevents complex dissociation. Formation of the closed conformation would also block the entrance of pyrophosphate and protect the 3'-dAMP from pyrophosphorolysis, as observed with the HCV RdRp (31). With 2'-C-Me-AMP misincorporation, the presence of UTP increased the steady-state rate, implying that UTP reduced the stability of the RdRp–RNA complex (Fig. 8). However, this result also indicates that UTP is still able to bind and form the RdRp–ssAU(2’CA)–UTP complex to exert such an effect. In contrast, addition of the noncognate nucleotide CTP had no effect on the steady-state rate (Fig. 8), likely because CTP would not bind sufficiently to form a ternary complex. RNA incorporated with 2'-C-Me nucleotide prevents the RdRp from forming the closed conformation and likely compromises the ability of the next incoming NTP to protect the 2'-C-Me nucleotide from pyrophosphorolytic excision, as observed with HCV RdRp (31).

Discussion

2'-C-Me ribonucleotides have become a very successful class of antiviral agents in the clinic, exemplified by the anti-HCV drug sofosbuvir (14). However, the molecular mechanism by which they terminate RNA synthesis has remained poorly understood. We chose PV RdRp (Fig. 1) as a model system, considering the vast amount of kinetic and structural data available, including our previous solution-state NMR studies (27, 35). PV RdRp can incorporate 2'-C-Me-AMP into RNA, which then leads to RNA chain termination (Fig. 2). We had previously established [methyl-13C]Met probes that report on RNA and NTP binding, including structural/dynamic changes representative of the less active open and more active closed conformations (Figs. 3 and 4). The NMR studies indicated that bind-
NMR studies of 2’-C-methylated nucleotide

Figure 7. Incorporation of 2’-C-Me-AMP prevents active-site closure upon binding the next incoming nucleotide. A, experimental design. PV RdRp (250 μM) is first incubated with 500 μM duplex RNA (ssAU or ssUU) and 4 mM nucleotide analogs, which, when incorporated, terminate RNA synthesis. The nucleotide analogs include the obligate chain terminators 3’-dATP, 3’-dGTP, and 2’-3’-ddATP and the nonobligate chain terminator 2’-C-Me-ATP (2’-CATP). Excess nucleotide analog was later removed through a desalting column, followed by addition of the second NTP (4–12 mM) to generate the RdRp–RNA–NTP ternary complexes. The D2O-based buffer consisted of 10 mM HEPES (pH 8.0), 200 mM NaCl, 0.02% NaN3, 5 mM MgCl2, and 10 μM ZnCl2. B–G, 1H-13C HSQC NMR spectra of different RdRp–RNA–NTP ternary complexes, including the RdRp–ssAU(3’-dA)–UTP (A), RdRp–ssAU(3’-dG)–UTP (B), RdRp–ssAU(2’-dA)–CATP (C), and RdRp–ssUU(2’-ddA)–CATP (D), RdRp–ssUU(2’-dA)–CATP (E), and RdRp–ssUU(2’-ddA)–CATP (F) complexes. Designations for the chain-terminated RNA can be found in the legend for Fig. 6, including the ssAU(2’CA) and ssUU(2’CA) RNA, which are ssAU and ssUU RNA with incorporated 2’-C-Me-AMP. Resonances belonging to the e-13CH3 groups of Met-6, Met-74, Met-187, Met-225, Met-354, and Met-394 are highlighted. NMR spectra were collected at 293 K using a Bruker Avance III 600 MHz spectrometer.

Binding and recognition of the 2’-hydroxyl group of the incoming ribonucleotide. It has been suggested that this amino acid substitution may interfere with initial 2’-C-Me-NT binding because of the steric clash from the additional Thr side-chain methyl group (44). However, it is also worth noting that the resistance variant has some ability to extend RNA synthesis past the incorporated nucleotide analog (13). Moreover, the S282T polymerase has lower catalytic efficiency and is associated with a fitness deficit (44, 45). These findings suggest that the S282T substitution may also have an effect on the open-to-closed conformational change necessary to incorporate the next incoming NTP, which would affect both polymerase efficiency and the ability to extend past the incorporated nucleotide analog. Additional NMR studies of the corresponding PV RdRp variant may provide further insight.

Other nonobligate chain terminators may likewise prevent the RdRp from fluctuating into the closed conformation. Nucleoside analogs may also interfere with other events of nucleic acid synthesis. For example, it has been shown that incorporation of the antiviral entecavir by the HIV reverse transcriptase results in “delayed” termination when the polymerase reaches the third nucleotide position downstream of the incorporated analog (46). Incorporation of the Thr–1106 analog results in backtracking of PV RdRp, which likely interferes with processivity and RNA yield (47). NMR studies of these analogs may provide new insights into molecular mechanisms behind translocation and processivity and provide new NMR signa-
was from Cambridge Isotope Laboratories. HisPur nickel-nitri- lotriacetic acid resin was from Thermo Scientific. Q-Sepharose Fast Flow was from GE Healthcare. All other reagents were of the highest grade, available from Sigma or Fisher.

**Production and purification of PV RdRp from heterologous expression in Escherichia coli**

PV RdRp was expressed in *E. coli* B834(DE3) pRARE cells using autoinduction and then purified as described previously (35, 48, 49). Protein samples used in NMR experiments were $^{13}$C-labeled by addition of [methyl-$^{13}$C] Met to the bacterial growth medium.

**NMR sample preparation and spectroscopy**

NMR samples were prepared as described previously (27, 35) with NMR buffer consisting of 10 mM HEPES (pH 7.5), 200 mM NaCl, 0.02% NaN$_3$, 5 mM MgCl$_2$, and 10 mM ZnCl$_2$. For generation of the RdRp–RNA binary complexes, PV RdRp (250 µM) was incubated with ssAU or ssUU RNA (500 µM duplex RNA) and 4 mM chain-terminating nucleoside triphosphate for 3–4 h before passage across a Zeba desalting column. The second NTP (4–12 µM) was then added to generate the RdRp–RNA ternary complexes. NMR experiments were performed on a Bruker Avance III 600 MHz spectrometer equipped with a 5-mm “inverse detection” triple-resonance ($^{1}$H/$^{13}$C/$^{15}$N) single-axis gradient TCI cryoprobe. $^{1}$H/$^{13}$C heteronuclear single quantum coherence (HSQC) spectra were generally acquired as 64 ($t_1$) × 512 ($t_2$) complex matrix, with 64–128 scans/increment and 1.0-s recovery delay at a temperature of 293 K (35).

**PV RdRp assays**

Steady-state kinetics assays were performed and analyzed as described previously (32). Reactions contained 50 mM HEPES (pH 7.5), 10 mM β-mercaptoethanol, 5 mM MgCl$_2$, 60 µM ZnCl$_2$, 1 μM PV RdRp, 20 µM ssAU RNA (i.e. 10 µM duplex RNA), and 500 µM appropriate NTPs. Reactions were initiated with addition of NTP(s), incubated at 30 °C, and then quenched at appropriate times by addition of EDTA to a final concentration of 50 mM. Products were analyzed by denaturing PAGE, and gels were visualized through use of a Phosphor Imager and quantified by using ImageQuant TL software (GE Healthcare).

**Inhibition of PV replication by 2′-C-methyladenosine**

The antiviral activity of 2′-C-methyladenosine was measured using procedures outlined previously (50). Infection with PV employed HeLa S3 host cells ($1 \times 10^5$) plated 1 day prior to treatment in 24-well plates. Cells were pretreated by addition of 2′-C-methyladenosine nucleoside at the specified concentration in fresh medium adjusted to a final concentration of 1% DMSO. After a 1-h incubation at 37 °C, the medium was removed, and cells were infected with PV ($1 \times 10^6$ pfu). Plates were incubated for 15 min at 23 °C, PBS was removed by aspiration, and fresh prewarmed (37 °C) medium containing the specified amount of nucleoside was added. The infection was allowed to proceed at 37 °C for 6 h. Cells were washed with PBS and collected after treatment with trypsin. Cells were pelleted by centrifugation, resuspended in PBS, and subjected to three freeze–thaw cycles. Cell debris was removed by centrifugation,
NMR studies of 2′-C-methylated nucleotide

and the supernatant containing the cell-associated virus was saved. Titer was determined by applying serial dilutions of supernatant to HeLa S3 monolayers (plated in 6-well plates 1 day before at 5 × 10^5 cells/well) and overlaying with growth medium containing low-melting-point agarose (1%). Plates were incubated for 2 days at 37°C. Then the agar was removed, and plaques were visualized by staining with crystal violet (1%) in aqueous ethanol (20%).

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References

1. Wang, L. S., D’Souza, L. S., and Jacobson, I. M. (2016) Hepatitis C: a clinical review. J. Med. Virol. 88, 1844–1855 CrossRef Medline
2. Sun, J., Hu, X. Y., and Yu, X. F. (2019) Current understanding of human enterovirus D68. Viruses 11, E9490 Medline
3. Yeaman, R., and Eikelboom, J. W. (2004) One hundred years of poliovirus pathogenesis. Virology 341, 9–16 CrossRef Medline
5. Collins, M. H., and Metz, S. W. (2017) Progress and works in progress: update on flavivirus vaccine development. Clin. Ther. 39, 1519–1536 CrossRef Medline
8. D’Ambrosio, R., Degasperi, E., Colombo, M., and Aghemo, A. (2017) Direct-acting antivirals: the endgame for hepatitis C? Curr. Opin. Virol. 24, 31–37 CrossRef Medline
9. Deval, J., Symons, J. A., and Beigelman, L. (2014) Inhibition of viral RNA polymerases by nucleoside and nucleotide analogs: therapeutic applications against positive-strand RNA viruses beyond hepatitis C virus. Curr. Topics Virol. 9, 1–7 CrossRef Medline
10. Carroll, S. S., Tomassini, J. E., Bosserman, M., Getty, K., Stahlhuber, M. W., Eldrup, A. B., Hall, D., Simoes, A. C., LaFemina, R., Rutkowski, C. A., Wolanski, B., Yang, Z., Migliaccio, G., De Francesko, R., et al. (2003) Inhibition of hepatitis C virus RNA replication by 2′-modified nucleoside analogs. J. Biol. Chem. 278, 11979–11984 CrossRef Medline
13. Migliaccio, G., Tomassini, J. E., Carroll, S. S., Tomei, L., Altamura, S., Bhat, B., Bartholomew, L., Bosserman, M. R., Ceccacci, A., Colwell, L. F., Cortese, R., De Francesko, R., Eldrup, A. B., Getty, K. L., Hou, X. S., et al. (2003) Characterization of resistance to non-obligate chain-terminating ribonucleoside analogs that inhibit hepatitis C virus replication in vitro. J. Biol. Chem. 278, 49164–49170 CrossRef Medline
14. Keating, G. M., and Vaidya, A. (2014) Sofosbuvir: first global approval. Drugs 74, 273–282 CrossRef Medline
15. Fung, A., Jin, Z., Dyatkina, N., Wang, G., Beigelman, L., and Deval, J. (2014) Efficiency of incorporation and chain termination determines the inhibition potency of 2′-modified nucleotide analogs against hepatitis C virus polymerase. Antimicrob. Agents Chemother. 58, 3636–3645 CrossRef Medline
16. Jin, Z., Tucker, K., Jin, X., Kao, C. C., Shaw, K., Tan, H., Symons, J., Behera, I., Rajwanshi, V. K., Dyatkina, N., Wang, G., Beigelman, L., and Deval, J. (2015) Biochemical evaluation of the inhibition properties of favipiravir and 2′-C-methyl-cytidine triphosphates against human and mouse norovirus RNA polymerases. Antimicrob. Agents Chemother. 59, 7540–7546 CrossRef Medline
17. Hansen, J. L., Long, A. M., and Schultz, S. C. (1997) Structure of the RNA-dependent RNA polymerase of poliovirus. Structure 5, 1109–1122 CrossRef Medline
18. Ferrero, D., Ferrer-Orta, C., and Verdaguer, N. (2018) Viral RNA-dependent RNA polymerases: a structural overview. Subcell. Biochem. 88, 39–71 CrossRef Medline
19. Appleby, T. C., Perry, J. K., Murakami, E., Barauskas, O., Feng, J., Cho, A., Fox, D., 3rd, Wetmore, D. R., McGrath, M. E., Ray, A. S., Sofia, M. J., Swaminathan, S., and Edwards, T. E. (2015) Viral replication: structural basis for RNA replication by the hepatitis C virus polymerase. Science 347, 771–775 CrossRef Medline
20. Gong, P., Cortus, M. G., Nix, J. C., Davis, R. E., and Peersen, O. B. (2013) Structures of coxsackievirus, rhinovirus, and poliovirus polymerase elongation complexes solved by engineering RNA mediated crystal contacts. PLoS ONE 8, e60272 CrossRef Medline
21. Gong, P., and Peersen, O. B. (2010) Structural basis for active site closure by the poliovirus RNA-dependent RNA polymerase. Proc. Natl. Acad. Sci. U.S.A. 107, 22505–22510 CrossRef Medline
22. Shu, B., and Gong, P. (2016) Structural basis of viral RNA-dependent RNA polymerase catalysis and translocation. Proc. Natl. Acad. Sci. U.S.A. 113, e4005–e4014 CrossRef Medline
23. Gohara, D. W., Crotty, S., Arnold, J. J., Yoder, J. D., Andino, R., and Cameron, C. E. (2000) Poliovirus RNA-dependent RNA polymerase (3Dpol): structural, biochemical, and biological analysis of conserved structural motifs A and B. J. Biol. Chem. 275, 25523–25532 CrossRef Medline
24. Steitz, T. A. (1993) DNA-dependent and RNA-dependent DNA-polymerases. Curr. Opin. Struct. Biol. 3, 51–38 CrossRef
25. Castro, C., Smidansky, E., Maksimchuk, K. R., Arnold, J. J., Korneeva, V. S., Götte, M., Konigsberg, W., and Cameron, C. E. (2007) Two proton transfers in the transition state for nucleotidyl transfer catalyzed by RNA- and DNA-dependent RNA and DNA polymerases. Proc. Natl. Acad. Sci. U.S.A. 104, 4267–4272 CrossRef Medline
26. Castro, C., Smidansky, E. D., Arnold, J. J., Maksimchuk, K. R., Moutha, L., Uchida, A., Götte, M., Konigsberg, W., and Cameron, C. E. (2009) Nucleic acid polymerases use a general acid for nucleotidyl transfer. Nat. Struct. Mol. Biol. 16, 212–218 CrossRef Medline
27. Yang, X., Smidansky, E. D., Maksimchuk, K. R., Lum, D., Welch, J. L., Arnold, J. J., Cameron, C. E., and Boehr, D. D. (2012) Motif D of viral RNA-dependent RNA polymerases determines efficiency and fidelity of nucleotide addition. Structure 20, 1519–1527 CrossRef Medline
28. Shu, B., and Gong, P. (2017) The uncoupling of catalysis and translocation in the viral RNA-dependent RNA polymerase. RNA Biol. 14, 1314–1319 CrossRef Medline
29. Sholders, A. J., and Peersen, O. B. (2014) Distinct conformations of a putative translocation element in poliovirus polymerase. J. Mol. Biol. 426, 1407–1419 CrossRef Medline
30. Garriga, D., Ferrer-Orta, C., Querol-Audí, J., Oliva, B., and Verdaguer, N. (2013) Role of motif B loop in allosteric regulation of RNA-dependent
RNA polymerization activity. J. Mol. Biol. 425, 2279–2287 CrossRef Medline
31. Deval, J., Powdrill, M. H., D’Abramo, C. M., Cellai, L., and Götte, M. (2007) Pyrophosphorolytic excision of nonobligate chain terminators by hepatitis C virus NS5B polymerase. Antimicrob. Agents Chemother. 51, 2920–2928 CrossRef Medline
32. Arnold, J. J., and Cameron, C. E. (2000) Poliovirus RNA-dependent RNA polymerase (3D(pol)). Assembly of stable, elongation-competent complexes by using a symmetrical primer-template substrate (sym/sub). J. Biol. Chem. 275, 5329–5336 CrossRef Medline
33. Liu, X., Musser, D. M., Lee, C. A., Yang, X., Arnold, J. J., Cameron, C. E., and Boehr, D. D. (2015) Nuclease but not sugar fidelity is maintained in the Sabin 1 RNA-dependent RNA polymerase. Viruses 7, 5571–5586 CrossRef Medline
34. Rosenzweig, R., and Kay, L. E. (2014) Bringing dynamic molecular machines into focus by methyl-TROSY NMR. Annu. Rev. Biochem. 83, 291–315 CrossRef Medline
35. Yang, X., Welch, J. L., Arnold, J. J., and Boehr, D. D. (2010) Long-range interaction networks in the fidelity and poliovirus RNA-dependent RNA polymerase studied by nuclear magnetic resonance. Biochemistry 49, 9361–9371 CrossRef Medline
36. Moustafa, I. M., Korboukh, V. K., Arnold, J. J., Smidansky, E. D., Marcotte, L. L., Gohara, D. W., Yang, X., Sánchez-Farrán, M. A., Filman, D., Maranas, J. K., Boehr, D. D., Hogle, J. M., Colina, C. M., and Cameron, C. E. (2014) Structural dynamics as a contributor to error-prone replication by an RNA-dependent RNA polymerase. J. Biol. Chem. 289, 36229–36248 CrossRef Medline
37. Moustafa, I. M., Shen, H., Morton, B., Colina, C. M., and Cameron, C. E. (2011) Molecular dynamics simulations of viral RNA-dependent RNA polymerases link conserved and correlated motions of functional elements to fidelity. J. Mol. Biol. 410, 159–181 CrossRef Medline
38. Yang, X., Liu, X., Musser, D. M., Moustafa, I. M., Arnold, J. J., Cameron, C. E., and Boehr, D. D. (2017) Triphosphate reorientation of the incoming nucleotide as a fidelity checkpoint in viral RNA-dependent RNA polymerases. J. Biol. Chem. 292, 3810–3826 CrossRef Medline
39. Gnädig, N. F., Beaucourt, S., Campagnola, G., Bordería, A. V., Sanz-Ramos, M., Gong, P., Blanc, H., Peersen, O. B., and Vignuzzi, M. (2012) Coxsackievirus B3 mutator strains are attenuated in vivo. Proc. Natl. Acad. Sci. U.S.A. 109, E2294–2303 CrossRef Medline
40. Korboukh, V. K., Lee, C. A., Acevedo, A., Vignuzzi, M., Xiao, Y., Arnold, J. J., Hemperly, S., Graci, J. D., August, A., Andino, R., and Cameron, C. E. (2014) RNA virus population diversity, an optimum for maximal fitness and virulence. J. Biol. Chem. 289, 29531–29544 CrossRef Medline
41. Freistadt, M. S., Vaccaro, J. A., and Eberle, K. E. (2007) Biochemical characterization of the fidelity of poliovirus RNA-dependent RNA polymerase. Virol. J. 4, 44 CrossRef Medline
42. Arnold, J. J., and Cameron, C. E. (2004) Poliovirus RNA-dependent RNA polymerase (3Dpol): pre-steady-state kinetic analysis of ribonucleotide incorporation in the presence of Mg2+. Biochemistry 43, 5126–5137 CrossRef Medline
43. Hedskog, C., Chodavarapu, K., Ku, K. S., Xu, S., Martin, R., Miller, M. D., Mo, H., and Svarovskaia, E. (2015) Genotype- and subtype-independent full-genome sequencing assay for hepatitis C virus. J. Clin. Microbiol. 53, 2049–2059 CrossRef Medline
44. Dutartre, H., Bussetta, C., Boretto, J., and Canard, B. (2006) General catalytic deficiency of hepatitis C virus RNA polymerase with an S282T mutation and mutually exclusive resistance towards 2’-modified nucleotide analogues. Antimicrob. Agents Chemother. 50, 4161–4169 CrossRef Medline
45. Powdrill, M. H., Tchesnokov, E. P., Kozak, R. A., Russell, R. S., Martin, R., Svarovskaia, E. S., Mo, H., Kouyos, R. D., and Götte, M. (2011) Contribution of a mutational bias in hepatitis C virus replication to the genetic barrier in the development of drug resistance. Proc. Natl. Acad. Sci. U.S.A. 108, 20509–20513 CrossRef Medline
46. Tchesnokov, E. P., Obikhod, A., Schinazi, R. F., and Götte, M. (2008) Delayed chain termination protects the anti-hepatitis B virus drug entecavir from excision by HIV-1 reverse transcriptase. J. Biol. Chem. 283, 34218–34228 CrossRef Medline
47. Dulin, D., Arnold, J. J., van Laar, T., Oh, H. S., Lee, C., Perkins, A. L., Harki, D. A., Depken, M., Cameron, C. E., and Dekker, N. H. (2017) Signatures of nucleotide analog incorporation by an RNA-dependent RNA polymerase revealed using high-throughput magnetic tweezers. Cell Rep. 21, 1063–1076 CrossRef Medline
48. Arnold, J. J., Bernal, A., Uche, U., Sterner, D. E., Butt, T. R., Cameron, C. E., and Mattern, M. R. (2006) Small ubiquitin-like modifying protein isopeptidase assay based on poliovirus RNA polymerase activity. Anal. Biochem. 350, 214–221 CrossRef Medline
49. Gohara, D. W., Ha, C. S., Kumar, S., Ghosh, B., Arnold, J. J., Wisniewski, T. J., and Cameron, C. E. (1999) Production of “authentic” poliovirus RNA-dependent RNA polymerase (3D(pol)) by ubiquitin-protease-mediated cleavage in Escherichia coli. Protein Expr. Purif. 17, 128–138 CrossRef Medline
50. Harki, D. A., Graci, J. D., Galarraga, J. E., Chain, W. J., Cameron, C. E., and Peterson, B. R. (2006) Synthesis and antiviral activity of S-substituted cytidine analogues: identification of a potent inhibitor of viral RNA-dependent RNA polymerases. J. Med. Chem. 49, 6166–6169 CrossRef Medline
51. Liu, X., Yang, X., Lee, C. A., Moustafa, I. M., Smidansky, E. D., Lum, D., Arnold, J. J., Cameron, C. E., and Boehr, D. D. (2013) Vaccine-derived mutation in motif D of poliovirus RNA-dependent RNA polymerase lowers nucleotide incorporation fidelity. J. Biol. Chem. 288, 32753–32765 CrossRef Medline