The SARS-CoV nsp12 Polymerase Active Site Is Tuned for Large-Genome Replication

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ABSTRACT

Positive-strand RNA viruses replicate their genomes using virally encoded RNA-dependent RNA polymerases (RdRP) with a common active-site structure and closure mechanism upon which replication speed and fidelity can evolve to optimize virus fitness. Coronaviruses (CoV) form large multicomponent RNA replication-transcription complexes containing a core RNA synthesis machine made of the nsp12 RdRP protein with one nsp7 and two nsp8 proteins as essential subunits required for activity. We show that assembly of this complex can be accelerated 5-fold by preincubation of nsp12 with nsp8 and further optimized with the use of a novel nsp8L7 heterodimer fusion protein construct. Using rapid kinetics methods, we measure elongation rates of up to 260 nucleotides (nt)/s for the core replicase, a rate that is unusually fast for a viral polymerase. To address the origin of this fast rate, we examined the roles of two CoV-specific residues in the RdRP active site: Ala547, which replaces a conserved glutamate above the bound NTP, and Ser759, which mutates the palm domain GDD sequence to SDD. Our data show that Ala547 allows for a doubling of replication rate, but this comes at a fidelity cost that is mitigated by using a SDD sequence in the palm domain. Our biochemical data suggest that fixation of mutations in polymerase motifs F and C played a key role in nidovirus evolution by tuning replication rate and fidelity to accommodate their large genomes.

IMPLICATIONS Replicating large genomes represents a challenge for RNA viruses because fast RNA synthesis is needed to escape innate immunity defenses, but faster polymerases are inherently low-fidelity enzymes. Nonetheless, the coronaviruses replicate their ~30-kb genomes using the core polymerase structure and mechanism common to all positive-strand RNA viruses. The classic explanation for their success is that the large-genome nidoviruses have acquired an exonuclease-based repair system that compensates for the high polymerase mutation rate. In this work, we establish that the nidoviral polymerases themselves also play a key role in maintaining genome integrity via mutations at two key active-site residues that enable very fast replication rates while maintaining typical mutation rates. Our findings further demonstrate the evolutionary plasticity of the core polymerase platform by showing how it has adapted during the expansion from short-genome picornaviruses to long-genome nidoviruses.

KEYWORDS RdRP, coronavirus, fidelity, polymerase

Positive-strand RNA viruses replicate their genomes using a virally encoded RNA-dependent RNA polymerase (RdRP) to carry out both minus and plus sense RNA synthesis. For the coronaviruses, this functionality is found in the nsp12 protein, and unlike most positive-strand RNA viruses that have single subunit RdRPs, the CoV polymerase is only active when complexed with the viral nsp7 and nsp8 proteins (1, 2). The structure of the polymerase from the original severe acute respiratory syndrome coronavirus 1 (SARS-CoV-1) strain showed a surprising stoichiometry wherein one copy of nsp12 interacted with one nsp7/nsp8 heterodimer and a second nsp8 monomer (3),
and the more than 40 structures of SARS-CoV-2 replicase solved in the past 2 years have expanded on this core structure. Notably among these, the two nsp8 molecules contain long N-terminal helices that interact nonspecifically with a long product RNA duplex as "sliding pole" processivity factors (4) (Fig. 1A). These helices also provide binding sites for two nsp13 helicases (5), while additional surfaces of nsp12 provide binding sites for nsp9 and a nsp10/14 heterodimer that contains both exonuclease and methyltransferase domains (6, 7). A growing body of biochemistry and structural biology data point to two distinct experimental contexts for the nsp12 RdRP subunit. The first is in a nsp12-nsp7/8-nsp8 "core polymerase" complex that can carry out processive elongation in vitro and is often used for biochemistry and nucleoside analog incorporation studies. The second is in the context of a much larger viral replication-transcription complex (RTC) that adds the nsp13 helicase, nsp14 exonuclease, N7-methyltransferase, nsp15 endonuclease, and nsp16 2'-O-methyltransferase subunits alongside the small nsp9 and nsp10 accessory proteins that stabilize subunit conformations and may indirectly chaperone RTC assembly. The full RTC structure has not been solved yet, but the multitude of nsp12 and other CoV nsp structures are providing hints about its composition (8–10), and a credible model for a 2.5-MDa RTC with hexameric symmetry has been built by combining interactions identified in cryo-electron microscopy (cryo-EM) and crystal structures of proteins from both SARS-CoV viruses with computational docking approaches (11).
Despite having a very large multicomponent RTC, the nsp12 RdRP active site and core fold is very similar to that of the 52-kDa picornaviral 3D<sub>pol</sub> enzymes that represent the smallest viral RdRPs (Fig. 1B). This is not altogether surprising as the nidoviruses evolved from the picornavirus-like supergroup of positive-strand RNA viruses (12, 13). Structural studies have shown that positive-strand RNA virus polymerases utilize an active-site closure mechanism that is based on a subtle movement of conserved motif A in the palm domain (14, 15), and biochemical and infectious virus studies show this core structure to be a platform upon which viruses can readily evolve their replication speed and fidelity to optimize fitness (16). Notably, replication rate and fidelity have been shown to be inversely correlated, with higher replication rates having a significant fidelity cost in both biochemical experiments and infectious virus studies (17–20). The CoV polymerase shares this structural core with a default open conformation active site, and mutations in motif A can alter replication fidelity based on the development of 5-fluoro-uracil resistance in mouse hepatitis virus CoV (21).

The ≈30-kB SARS-CoV genome is three to four times larger than those of more common RNA viruses such as flaviviruses and picornaviruses, and this size presents two key biochemical challenges for the CoV replication machinery. First, it must be very fast in order to replicate the entire genome before innate immune responses are activated in cells, and second, it must retain sufficiently high fidelity so as to maintain genetic integrity. Current data show that the CoV core replicase elongates at 150 to 200 nucleotides (nt)/s (22–25), which is more than twice as fast as poliovirus 3D<sub>pol</sub>, and based on prior RdRP studies, this is likely to be accompanied by a fairly high mutation rate inherent to the polymerase active-site architecture (16). Several studies show nsp12 readily misincorporates native and modified nucleotides <i>in vitro</i>, including the nucleoside analogs remdesivir and favipiravir that show antiviral activity in cell culture and molnupiravir (N4-hydroxycytidine) that is approved as a CoV antiviral (24, 26, 27). Careful comparative measurements of nsp12 fidelity with other polymerases have not been made, and it is not known whether there are molecular features of nsp12 that allow it to maintain fidelity in light of its fast replication rate. The large-genome CoVs are also unique among RNA viruses in having an exonuclease-based repair mechanism via the ExoN domain of nsp14 that can offset the inherently reduced fidelity of a faster polymerase (28).

While the core of SARS-CoV nsp12 polymerase shares the canonical RdRP fold and mechanism, it does have two notable active-site differences from the typical positive-strand RNA virus polymerase (Fig. 1C); Ala547 within motif F reflects the loss of a key glutamic acid residue that normally sits above the active site and aids in positioning an arginine side chain and the NTP for catalysis and Ser759 within motif C results in a SDD sequence instead of a GDD sequence in the palm domain that interacts with the nascent RNA strand. Thus, positive-strand RNA virus polymerases generally use a glutamate-GDD configuration of these motifs F and C residues that we will denote as E:GDD, but CoVs have an A:SDD configuration (Fig. 1D). The SDD is present in all nidoviruses, but interestingly, the shorter genome arteriviruses have Q:SDD with a motif F glutamine that lacks the formal negative charge of the glutamate and would therefore have a weaker interaction with the active-site arginine.

In this study, we sought to determine how these two noncanonical features of the CoV nsp12 active site affect replication rate and fidelity. Detailed functional studies have been hampered by the difficulty of obtaining pure, highly active coronavirus replicase preparations. To overcome this complication, we first developed a new nsp8L7 fusion protein and used it to elucidate the pathway for core replicase assembly and then used assembled complexes in rapid kinetics and misincorporation studies to assess the biochemical effects associated with the Ala547 and Ser759 residues. Our data show that having Ala547 in motif F more than doubles the replication rate, and while this comes at a significant fidelity cost, the effect is mitigated by using SDD instead of GDD within motif C as a new fidelity control point. As a result, the overall nucleotide misincorporation level of nsp12 is on par with that of the much slower...
poliovirus polymerase. These compensatory biochemical effects in CoV nsp12 show a new link between the core RdRP active-site mechanisms and maximizing virus fitness and provide function-based insights into nidoviral polymerase evolution.

RESULTS
Design of a highly active SARS-CoV-2 core replicase. Examination of the cryo-EM structure of the SARS-CoV-2 core replicase reveals that the nsp8 C terminus is in the vicinity of the nsp7 N terminus (3, 8). We designed nsp8L7, a single-chain version of the nsp7/8 heterodimer, with the goal of expediting the kinetics and specificity of complex formation. The nsp8L7 protein, in which nsp8 is connected to the N terminus of nsp7 via a 6-residue GSGSGS linker, allows us to delineate the biochemical roles of the two distinct nsp8 molecules in the replicase complex while retaining the long nsp8 N-terminal α-helix that interacts with the product RNA duplex. All proteins used in this study were expressed in Escherichia coli using T7 promoters and purified through metal-affinity, ion-exchange, and size-exclusion chromatography (Fig. 2A). Purification yield of nsp12 was fairly low at ≈1 mg/liter of bacterial culture, and it copurified through metal-affinity chromatography with a number of bacterial proteins (29) that could be largely removed via a shallow salt gradient on a high-resolution MonoQ anion-exchange column (Fig. 2A). The size-exclusion data showed nsp12 (109 kDa), nsp8 (24 kDa), and nsp7 (11 kDa) eluting near their expected positions based on

![FIG 2](image)

Protein purification and RNA binding. (A) SDS-PAGE of purified SARS-CoV-2 nonstructural proteins, including an engineered nsp8L7 fusion protein with a 6-residue linker (8L7* contains an 18-residue linker); MonoQ anion-exchange column chromatogram, and SDS-PAGE gel showing separation of nsp12 into “pure” (P) and “less pure” (LP) sample pools as indicated on the gel with brackets. (B) Superposed size-exclusion chromatograms showing that nsp12 (109 kDa), nsp8 (24 kDa), and nsp7 (11 kDa) elute at their expected sizes, while nsp8L7 (33 kDa) is well shifted from the expected elution volume. (C) Fluorescence polarization (FP) RNA-binding data for each of the nsp constructs obtained in 50 mM NaCl, pH 7.
standards, but the nsp8L7 (33 kDa) fusion protein eluted at an apparent molecular weight of ≈100 kDa (Fig. 2B). This indicates nsp8L7 forms oligomers in solution, which is consistent with several crystal structures showing that the nsp7/nsp8 heterodimer can itself form (nsp7/8), dimers, and these dimers may further oligomerize into ((nsp7/nsp8))₂ tetramers and larger hexadecamers via interactions involving the long N-terminal helix of nsp8 (30–33). RNA-binding experiments in the presence of 50 mM monovalent salt show that nsp8, nsp8L7, and nsp12 all bound a short hairpin primer-template RNA with ≈200 nM $K_d$ values, while nsp7 binds weakly with a 7 μM $K_d$ (Fig. 2C). Deletion of the first 59 residues of the nsp8L7 fusion protein reduced binding affinity more than 80-fold to ≥14 μM.

**Core replicase assembly.** Assembly of the core replicase complex, here defined as nsp12-nsp7/8-nsp8, from individual components and a hairpin primer-template RNA (Fig. 3A) was observed to be both slow and inefficient at micromolar concentrations, requiring ≈15 min and elongating only half of the RNA hairpin primer provided (Fig. 3B). Final concentrations were 1 μM RNA and 2.5 μM nsp12 either with 5 μM nsp7 and 10 μM nsp8 or with 5 μM nsp8 and 5 μM nsp8L7. This result is based on using short 15-s burst-phase elongation reactions to instantaneously assay the amount of competent complex present after various assembly times, and the data therefore do not reflect multiple turnover product formation over the course of the entire assembly window. Assembly with nsp8L7 was ≈2-fold faster and ≈35% more efficient than assembly with individual nsp7 and nsp8 proteins (Fig. 3B and D). Preincubation of nsp12 with nsp8 prior to the addition of nsp8L7 and RNA accelerated the assembly of functional replicase ≈3-fold, but preincubation of nsp12 with nsp8L7 did not (Fig. 3C and D), indicating that the slow assembly from separate components is due to the lone nsp8 binding to the nsp12 pinky finger domain. Overall, the combination of nsp8 preincubation and using the nsp8L7 fusion instead of nsp7+nsp8 led to a 7.5-fold increase in core replicase assembly efficiency. Preassembly of nsp12+nsp8 followed by nsp8L7+RNA addition was used for all subsequent experiments unless otherwise stated.

To assess the lifetime of the nsp12-8 heterodimer, we assembled the complex, purified it by size-exclusion chromatography, and then made a series of reinjections of the purified sample on the same size-exclusion column over an 18-h period. These data showed that the heterodimer dissociated to a new equilibrium with a time constant of ≈6 h at room temperature in 100 mM NaCl, pH 7, 10% glycerol (Fig. 3E), and the final concentrations based on measured peak areas were ≈265 nM complex and ≈640 nM free nsp12 and nsp8, yielding an estimated $K_d$ of ≈1.5 μM for the nsp12-nsp8 interaction. We also note that the amount of actively elongating complex formed varied somewhat among different nsp12 preparations and sample pools (Fig. 2A), and we attribute this to conformational heterogeneity of nsp12 and the presence of copurifying bacterial proteins and chaperones that primarily interfere with formation of the nsp12-nsp8 heterodimer. The final assembly step after addition of nsp8L7 and RNA was consistently 2 to 3 min, and regardless of overall assembly efficiency, the active complexes formed exhibited consistent elongation rates and misincorporation levels in the assays described below.

**Processive elongation rates.** Processive elongation rates and nucleotide $K_{m,app}$ values were determined using a fluorescence based stopped-flow assay to measure the time needed to replicate a 26-nucleotide-long single-stranded RNA template (Fig. 4A). The assay is based on a fluorescence signal change triggered by the polymerase approaching the very 5’ end of the template strand RNA and thus reports only on the population of elongation competent replicases (34). Notably, the assay result is insensitive to the replicase assembly efficiency as the signal from nonelongated RNAs remains constant; it only requires enough competent complexes to generate a fluorescence change with sufficient signal-to-noise ratio to be interpretable. Using titrations with an equimolar mixture of the four NTPs, we examined the concentration dependence of processive elongation rates as a function of temperature at pH 6.5 and as a function of...
pH at 30°C (Fig. 4B). The extrapolated maximal rates observed across these conditions ranged from 23 ± 6 to 155 ± 15 nt/s with NTP \( K_m \),app values in the 50 to 100 \( \mu \)M range.

Based on these data, we carried out an expanded temperature range and pH study at a single 350 \( \mu \)M NTP concentration, i.e., at near-maximal elongation rate conditions. The resulting data showed the CoV replicase to be an extremely fast RdRP, with processive elongation rates of 250 to 300 nt/s at 37°C and pH 7.5 (Fig. 4C). Individual deletions of the nsp8 N-terminal helix regions that do not directly contact nsp12, i.e., nsp8-D75 and nsp8L7-D59 (Fig. 1A), did not significantly affect the processive elongation rate. These deletions did reduce the amount of competent replicase formed, which led to lower signal-to-noise ratio in the stopped-flow data, but reliable elongation rates

**Fig 3** Core CoV replicase assembly. (A) Hairpin RNA (HP) used to assay replicase assembly via 15-s elongation reactions yielding a T GAGA (+4, gray) product imaged via a 5' IRDye800 label. (B) 7 M urea/Tris-borate-EDTA (TBE)-PAGE showing that the 8L7 fusion protein improves both the rate and efficiency of complex assembly compared to individual nsp7 and nsp8 subunits in reactions initiated by mixing all the components. (C) A 30-min preincubation of nsp12 with nsp8, but not nsp8L7, accelerates the core replicase assembly process. (D) Quantification of assembly times obtained by fitting the fraction of RNA elongated to a single exponential curve. The data in panels B to D reflect the final assembly step done with 1 \( \mu \)M RNA, 2.5 \( \mu \)M nsp12, and either 5 \( \mu \)M nsp7 or 5 \( \mu \)M nsp8L7. The 30-min preincubation steps were done at 2°C final concentrations, i.e., 5 \( \mu \)M nsp12 and 10 \( \mu \)M cofactor proteins. (E) Size-exclusion chromatography was used to monitor the temporal stability of the nsp12/8 complex at room temperature. Area measurements of the nsp12/8 and nsp12 peaks showed the complex dissociates in \( \approx 6 \) h (inset) and has a \( \approx 1.5 \) \( \mu \)M \( K_d \) based on estimated equilibrium concentrations.
could still be determined (Fig. 4D). However, we could not determine the elongation rate of the double deletion combination, nsp8-Δ75 with nsp8L7-Δ59, because it did not form enough active complex.

**Replication fidelity.** Using the burst-phase assay format with a self-priming hairpin RNA, we observed significant amounts of misincorporation products indicating that the wild-type CoV replicase complex is also a low-fidelity RdRP. With a template whose cognate elongation product would be AACUAGACAG, we observed the expected elongation product (AA) when only ATP (50 μM) was provided and the expected product (AAC) when both 50 μM ATP and low 5 μM concentrations of CTP were provided (Fig. 5A). However, when the CTP concentration was increased to a mere 50 μM, we observe CTP:A mismatch incorporation, using a nomenclature that reflects a CTP paired opposite a template strand adenosine, and this generated a product that was further elongated to an AACcA (Fig. 5A). The misincorporation efficiency was CTP concentration dependent, and at 400 μM CTP we observed ~50% of the initiated material being elongated past the +3 species in the 15-s burst reactions (Fig. 5B). Essentially identical misincorporation levels were observed using the nsp8L7 fusion protein or the

FIG 4 Stopped-flow fluorescence elongation assays. (A) Primer-template RNA construct composed of a primer strand (blue) and two-piece template strand (red, gray) that yield a long 29-bp priming duplex and 26-nt single-stranded template with a fluorescein (FL) at the distal 5′ end of the template. (B) Stopped-flow fluorescence data showing the NTP-dependent shortening of the initial lag phase that reflects processive elongation before the nsp12 complex arrives at the 5′ end and triggers an increase in the fluorescein signal. NTP concentration dependence curves were fit to determine maximal elongation rates and Km,app values for eight full titrations carried out at various temperatures and pH values. (C) Temperature and pH dependence of near-maximal elongation rates obtained at a single 350 μM NTP concentration. (D) Stopped-flow analysis of elongation rates using the wild-type nsp8 and nsp7 cofactors and N-terminal truncations of nsp8 and nsp8L7 (see Fig. 1A for truncation sites). The N-terminal truncations destabilize the elongation complex and reduce signal amplitudes because fewer active complexes are formed, but lag phases can be reliably measured as demonstrated by scaling up and 5-point smoothing of their highest NTP concentration data traces (red to gray).
individual nsp7 and nsp8 subunits (Fig. 5B). With the increasing amounts of +5 product generated at higher CTP concentrations, we also observe a second misincorporation that we attribute to an ATP:C mismatch, i.e., adenosine in place of guanosine, to produce a species that is further elongated to an AACcAaACA product (Fig. 5A).

We did not observe the terminal ATP:C mismatch that would generate the full-length product and attribute this to the instability of the elongation complex when there are no downstream nucleotides remaining in the nsp12 template entry channel. We quantitated the level of misincorporation using three gel bands: the band reflecting initiation and cognate AAC addition, the band that requires a CTP:A mismatch at the position, and the band that requires an ATP:C mismatch at the position. At the molecular level, these misincorporation reactions occur in the context of processive elongation because each replicase complex begins with the original hairpin primer, and any intermediate +3 or +5 RNA products that dissociate prior to mismatch incorporation are unlikely to reform new elongation complexes within the 15-s reaction time. Using a similar self-priming RNA hairpin in which the cognate product was GAGACCAUUC, we observed only the cognate +10 product and this forms a very stable hairpin that does not denature on the gels and therefore migrates faster than the misincorporation product RNAs.

We then compared these nsp12 misincorporation levels with those obtained from the picornaviral poliovirus and coxsackievirus B3 3Dpol enzymes. For a direct comparison with the nsp12 data, these experiments were done by preincubating 3Dpol and hairpin RNA and then adding the required nucleotides for processive elongation and

![Diagram of RNA hairpin and gel bands](image)
mismatch incorporation; unlike in most of our prior studies, we did not preincubate with the first nucleotide(s) to generate stalled elongation complexes prior to elongation. Both 3Dpol enzymes also showed CTP concentration-dependent CTP:A misincorporation leading to the +5 product, and the subsequent ATP:C misincorporation leading to the +9 product (Fig. 5A). CV 3Dpol showed higher fidelity than PV 3Dpol (Fig. 5B), which is consistent with prior comparisons of GTP:U mismatch incorporation by the picornaviral enzymes (16). One particularly interesting observation from these data is that the amount of mismatch products made by nsp12 and poliovirus 3Dpol are roughly comparable, especially at lower CTP concentrations (Fig. 5B), suggesting that there are features of the nsp12 active site that maintain fidelity despite its significantly faster elongation rate.

**Comparison with a canonical RdRP active site.** To address the origin of this fidelity compensation, we generated individual A547E and S759G mutations in nsp12, and conversely E161A and G327S mutations in poliovirus 3Dpol (Fig. 1C and D), and analyzed their effects on processive elongation with rapid stopped-flow kinetics and on mismatch incorporation with burst-phase assays (Fig. 6). Introducing the motif F glutamic acid into the nsp12 active site reduced the maximal elongation rate 2.4-fold from 105 to 44 nt/s, and the SDD-to-GDD mutation reduced rate by a greater 7.5-fold factor to only 14 nt/s, and the double mutation reduced it 15-fold to 7 nt/s (Fig. 6A). These mutations also decreased the NTP $K_{\text{m,app}}$ values 2- to 3-fold, from 100 $\mu$M to 59, 29, and 26 $\mu$M. Fidelity increased similarly with both the A547E mutation and double mutations based on reductions in both CTP:A and ATP:C mismatches and decreased with S759G based on significantly higher levels of CTP:A mismatch incorporation (Fig. 6B). The plot in Fig. 6B reflects CTP:A mismatch events, and an analysis of only the ATP:C mismatches, i.e., +9 band divided by sum of +5 and +9 bands, showed similar effects.
with 26% ± 2% ATP:C product for wild-type nsp12 that decreased to 11 ± 1% with the higher fidelity A547E mutation, increased to 40 ± 3% with the lower fidelity S759G mutation, and decreased to 3 ± 1% with the double mutation. Thus, the two noncanonical residues in the native nsp12 active site have complementary functions that both benefit large-genome synthesis, with Ala547 accelerating replication rate and Ser759 improving fidelity.

In the context of poliovirus 3Dpol, an E161A mutation in motif F increased the processive elongation rate 1.6-fold and increased $K_{\text{m,app}}$ (Fig. 6A), effects that were opposite of those observed with the reverse mutation in nsp12, and again showing that the smaller alanine in motif F yields a faster polymerase. Interestingly, the E161A mutation also significantly destabilized the 3Dpol-RNA complex, resulting in lower signal amplitudes in the raw stopped-flow data (not shown) and inefficient initiation in the burst-phase elongation experiments to assess fidelity (Fig. 6B). The observed misincorporation level of the E161A mutant was comparable to that of wild-type 3Dpol, but note that this may be an underestimate of the true misincorporation efficiency because the unstable 3Dpol complexes may dissociate from the +3 product RNA faster than they misincorporate to generate the +4 and longer species. Last, the introduction of SDD into 3Dpol motif C via the G327S mutation slowed processive elongation 3.7-fold and significantly reduced misincorporation levels, with CTP:A being reduced 8-fold and ATP:C mismatch products being barely detectable. This is a clear increase in fidelity that is consistent with the higher fidelity associated with SDD over GDD in nsp12. We also attempted assays with the double mutant 3Dpol, but this was even less stable than the E161A single mutant. We could not obtain any elongation signals in the stopped-flow assay and barely detected the cognate +3 product in the burst assays (Fig. 6B), precluding any misincorporation assessment.

Overall, our data show that the nsp12-nsp7/8-nsp8 core CoV replicase complex could be efficiently assembled by using a nsp8L7 fusion protein in place of the nsp7/8 heterodimer and preincubating nsp12 and nsp8. This complex is a very fast viral polymerase with elongation rates approaching 300 nt/s at 37°C, and active-site mutants show this fast rate is enabled by nsp12 having an atypical alanine in motif F alongside a SDD sequence within the palm domain motif C that helps maintain fidelity.

**DISCUSSION**

Virus mutation rates as generated by the molecular replication machinery and the related mutation frequencies as observed in the resulting virus populations are inherently difficult values to determine (35). They reflect a complicated interplay of polymerase fidelity, co- and postsynthesis repair mechanisms, fitness, selection and adaptation effects, RNA modifications by host factors, the biochemical environment of the host cell, and even the molecular methodologies used to detect mutations. Nonetheless, careful comparative analyses indicate that positive-strand RNA virus mutation rates are 200- to 2,000-fold higher than what is seen for DNA-based genome replication (36–38). This largely reflects the lack of proofreading and repair systems in most RNA viruses, and the inherent error rates of DNA polymerase and viral RdRPs are comparable (39). However, the coronaviruses exhibit $\approx$10-fold higher replication fidelity than most RNA viruses, and this is essential for the survival of these nonsegmented large-genome viruses. While much of this mutation rate reduction can be attributed to the ExoN proofreading enzyme that is unique to the large-genome nidoviruses, it is noteworthy that all nidoviral polymerases have two key active-site mutations that differentiate them from the RdRPs of smaller genome positive-strand RNA viruses. This raises the question of how the nidoviral RdRPs have adapted to enable higher replication rates and what effect this has on inherent polymerase fidelity. In this study, we address this issue by first elucidating a pathway for efficient *in vitro* assembly of the SARS-CoV-2 core replicase complex composed of nsp12-nsp7/8-nsp8 and then using these complexes to assay elongation rates and nucleotide mismatch incorporation levels.

**Core replicase assembly is rate limited by the nsp12-nsp8 interaction.** The assembly of functional CoV replicase complexes is inefficient, leading to seemingly low
elongation activity on a time scale of tens of minutes in assays in which all the protein, RNA, and NTP components are mixed simultaneously. High molar excess concentrations of nsp7 and nsp8 are often used in studies to increase efficiency, but even then, the assembly is not complete based on large amounts of unused primer. However, the subpopulation of RNA that is assembled into complexes is rapidly elongated in quench-flow and single-molecule experiments (22–24). These data underscore that CoV initiation is a multistep process that is at a minimum composed of slow assembly of the (nsp12-nsp7/8-nsp8-RNA) core replicase complex followed by rapid elongation. To separate these two confounding biochemical activities, we first sought to identify the slow step in the core replicase assembly process.

Building on the prior nsp7L8 fusion version of the nsp7/nsp8 heterodimer (1) and considering the now available structures of the CoV core replicase, we reversed the order of the two proteins to generate a single-chain nsp8L7 protein mimicking the nsp8/nsp7 dimer bound to nsp12. This greatly enhanced assembly of a highly active core replicase, and based on the order of addition data, we propose a model in which initial formation of a nsp12-nsp8 complex is the rate-limiting factor in replicase assembly that is followed by the rapid addition of nsp8L7 and RNA. Our finding that a nsp12-nsp8 complex forms first is consistent with the observation that cotranslational cleavage of a nsp7-nsp8-nsp12 fusion protein in insect cells yielded a purifiable nsp12-nsp8 complex that lacked nsp7 (2, 27). Importantly for in vitro experiments, using our two-step protocol with nsp12-nsp8 preassembly allows us to generate active replicase complexes at relatively low and close-to-stoichiometric concentrations of the nsp8 and nsp8L7 cofactor proteins. This minimizes formation of alternative complexes as these cofactors are themselves prone to oligomerization, as seen in our data (Fig. 2B) and characterized previously to yield complexes that may compete for RNA binding and elongate RNA independently of nsp12 (32, 33, 40–42).

**nsp12 rate and fidelity.** Stopped-flow rapid kinetics showed nsp12 elongation rates of almost 300 nt/s, making the CoV replicase the fastest viral RdRP characterized to date (Fig. 4C). These rates are consistent with the ≈170 nt/s observed at 37°C in magnetic tweezers studies (22, 25) and quench-flow experiments using both SARS-1 and SARS-2 replicases (23, 24). Notably, they are 3-fold faster than the ≈90 nt/s rates observed for poliovirus 3Dpol and 15-fold faster than the ≈20 nt/s coxsackievirus 3Dpol that share the same core structure and active-site closure mechanism (16, 34). The two nsp8 sliding pole helices do not significantly affect processive elongation rates (Fig. 4D), but their deletion reduced RNA binding (Fig. 2C) and significantly lowered amplitudes in the stopped-flow studies (Fig. 4D), consistent with the sliding poles acting as processivity factors that stabilize the elongation complex.

To determine whether there are coronavirus-specific features responsible for the remarkably high nsp12 RNA synthesis rate, we examined how the noncanonical Ala547 instead of a glutamate in motif F and Ser759 instead of a glycine in motif C affected rate and fidelity. Stopped-flow elongation data showed that both residues make significant contributions to the high replication rate of nsp12, with individual A547E and S759G mutations reducing elongation rates 2- and 7-fold, respectively. The two mutations had opposite effects on fidelity, which increased with the A547E mutation and decreased significantly with the S759G mutation (Fig. 6B). The S759G mutation provided a very clear indication of reduced fidelity in that it increased misincorporation levels almost 2-fold while reducing the processive elongation rate 7-fold, implying a ≈14-fold decrease in nucleotide specificity. These data reflect misincorporation during processive elongation because given the slow core replicase assembly rate, the 15-s reaction time in our burst assays is too short to allow for multiple turnover reinitiation via RNA release and rebinding.

We then converted a canonical RdRP active site into a CoV type active site by making the opposite mutations in poliovirus 3Dpol. Mutating the motif F glutamate to alanine (E161A) increased 3Dpol elongation rate 1.6-fold, which is consistent with rate reduction observed with the opposite A547E mutation in nsp12. The observed fidelity effect arising from E161A was minor (Fig. 6B), but note that this may reflect an
underestimate of the true fidelity because the mutation itself significantly destabilized the 3Dpol-RNA complex. The RNA dissociation rate may therefore be comparable to the misincorporation rate, which would reduce the amount of misincorporated product observed in these burst-phase assays. The GDD to SDD mutation in 3Dpol showed a \( \approx 3.7 \)-fold reduction in processive elongation rate and an \( \approx 8 \)-fold reduction is misincorporation levels, suggesting it is a higher fidelity polymerase. However, as both effects are rate reductions and the inferred increase in nucleotide specificity is only \( \approx 2 \)-fold, we cannot clearly establish from the burst-phase data if the reduced misincorporation level is a true fidelity effect or simply a reflection of slower kinetics.

**Polymerase adaptations for large-genome replication.** RNA virus genomes are limited by the tradeoffs inherent in maintaining genetic integrity in the context of replication by low-fidelity RNA-dependent RNA polymerases, and consequently the majority of RNA viruses have genomes in the 4- to 12-kb range. Larger genomes require more time for replication, but using a faster RdRP can be counterproductive because it tends to also increase mutation rates. Thus, the expansion of RNA genomes past the \( \approx 12 \)-kb threshold typified by most flaviviruses requires both an increase in replication rate and a decrease in mutation rate, the latter of which could be achieved by either improving RdRP fidelity or by acquiring error-correcting functionality (12, 13, 37). The nidoviruses have undergone such an expansion, resulting in genomes of up to 32 kb, the existence of which is largely attributed to the acquisition of an error-correcting exonuclease (ExoN) enzyme. However, ExoN is present only in large-genome (>20 kb) nidoviruses, i.e., the coronaviruses and toroviruses, while smaller genome arteriviruses and some flavivirus-like viruses have managed to expand their genomes to 13 to 16 kb and 27 kb, respectively, without an ExoN (37). This suggests that the inherent speed and fidelity of the core viral RdRP platform can be modulated to allow for some expansion of the viral genome without adding proofreading error correction.

At the structural level, the positive-strand RNA virus polymerases share a core fold and palm-domain active-site closure mechanism that is based on a subtle movement of conserved motif A (10, 14, 43, 44). The active site itself includes three aspartates that are essential for magnesium coordination and catalysis and therefore present in all replicative polymerases (45). Beyond this, the exact amino acid sequence determines elongation rate and fidelity, with particularly strong effects arising from substitutions within motifs A and D (16) and more subtle effects coming from extended protein dynamics networks within the fingers domain (46–48). This led to a model in which the palm domain-based active-site closure mechanism provides a structural framework upon which individual viruses can evolutionarily fine tune RdRP function to optimize fitness (16, 18, 49). Outside of the immediate catalytic center, viral RdRPs commonly retain important residues that trigger the closure mechanism or interact directly with the RNA and NTP substrates. Among the latter is the glycine in the GDD sequence of the motif C \( \beta \)-hairpin structure that is located below the priming nucleotide ribose and a glutamate-arginine salt bridge in motif F that is positioned directly above the bound NTP \( \alpha / \beta \)-phosphates (Fig. 1C). Interestingly, the nidoviral polymerases have different residues at both sites, leading us to propose a two-step process whereby the core RdRP structure and mechanism has adapted to support large-genome replication.

The molecular mechanism underpinning the effects of these mutations is best explained by first considering the catalytic geometry of picornaviral polymerases, in which high-resolution crystal structures have mapped out the entire RdRP nucleotide addition cycle (15). RNA-templated polymerases, i.e., RdRPs and reverse transcriptases, have a conserved arginine within motif F and the ring finger structure that is the structural homolog of an O-helix lysine found in DNA templated single subunit polymerases (14). Multiple structures show the arginine positioned directly above the NTP \( \alpha / \beta \)-phosphate linkage in precatalytic RdRP-RNA-NTP complexes (14, 50–52), where it imposes an electrostatic positioning constraint on the NTP \( \alpha \)-phosphate that will favor catalysis when a correctly base-paired NTP is bound in the active site (Fig. 1C). In the picornaviral RdRPs, this arginine (Arg174) is held in place by a salt bridge to a glutamate (Glu161) on the other \( \beta \)-strand of the ring finger, as shown in Fig. 1C.
and schematically illustrated in the E:GDD panel of Fig. 7. Notably, this glutamate-arginine guanidinium group interaction is well ordered in PV 3Dpol even in the absence of bound RNA and/or NTP (53). Removal of the salt bridge via the E161A mutation in poliovirus poly-

merase increases the elongation rate (Fig. 6A), and this likely arises from a combination of increased arginine flexibility, schematized by the halo around the R in the A:GDD panel of Fig. 7, and a steric load of the NTP into the active site; the rigid Glu-Arg salt bridge is also a steric barrier that the incoming NTP needs to physically bypass in order to base-pair with the templating nucleotide.

In this context, it is notable that all the nidoviral polymerases have two key active-site mutations: loss of the motif F glutamate and conversion of the palm domain motif C GDD sequence into an SDD. Comparing the 46 currently available nsp12 structures via the coro set of polymerase superpositions (9, 10) shows the motif F arginine (Arg555) adopting a wide range of positions in both the absence and presence of RNA and NTP (Fig. 1C). We propose that it is this inherent flexibility and lack of steric con-

straints associated with the Glu-Arg salt bridge that enables rapid NTP loading into the active site and fast elongation rates. The inherent molecular tradeoff of this fast elongation rate is lower fidelity due to fewer constraints on the NTP α-phosphate positioning during catalysis. To overcome this obstacle, the nidoviruses have established a new fidelity checkpoint by using a SDD instead of GDD sequence within motif C in the palm domain. The cryo-EM structures of several nsp12-RNA complexes show the SDD serine side-chain hydrogen bonding with the 2′ hydroxyl of the priming ribose (Fig. 1C) (54–59), leading us to propose that the mechanistic explanation for the improved fidelity of SDD is a reinforcement of correct priming ribose positioning in the
active site. In effect, using SDD in motif C shifts a fidelity checkpoint from NTP positioning to primer 3’ end positioning, and this allows for dramatic increases in elongation rate by releasing structural constraints on the motif F arginine. Conversely, having both fidelity control points present at the same time, as in the PV G327S mutant (Fig. 6B), results in a very high-fidelity enzyme, but also a slow enzyme, and consequently any virus carrying this combination would have significantly reduced growth rate. We speculate that this E:SDD configuration enzyme is slow because of conflicting interactions that independently constrain the positions of the NTP and the priming ribose into different sites, leading to a combined geometry that is less than ideal for catalysis (Fig. 7).

When considering the polymerase adaptations in going from short- to long-genome RNA viruses, it is also insightful to consider the intermediate length 13- to 16-kb arterivirus RdRPs. They have a Q:SDD configuration with a motif F glutamine that is a subtle mutation from the canonical glutamate and would likely interact similarly with the arginine, but the interaction would be weaker by virtue of being nonionic. The prediction of our model is that replication rate would increase, and this would then enable replication of slightly longer genomes in a time frame that can escape innate immune pressure, but fidelity would decrease because of the less restricted active-site geometry. We propose that it is this fidelity constraint that led to the subsequent emergence of the SDD sequence in motif C and shifting the fidelity checkpoint away from NTP α-phosphate positioning and toward priming ribose positioning. The presence of SDD in conjunction with a slight relaxation of the arginine interaction via a glutamine, e.g., Q:SDD, allowed for stable 13- and 16-kb arterivirus genomes. The jump to even larger genomes would require even faster polymerases, which could now be easily achieved by further loosening the constraints on the motif F arginine, resulting in the minimal alanine residue and A:SDD configuration found in the 26- and 32-kb coronaviruses and toroviruses or S:SDD found in roniviruses. However, at these very fast polymerase rates, the SDD fidelity checkpoint would no longer be sufficient for maintaining genome integrity, and therefore, these viruses also acquired ExoN domains to carry out postcatalysis error correction as a separate step. From the studies presented here, we cannot infer whether the arteriviral polymerases evolved from the picornaviral-like supergroup in parallel with the large-genome nidoviruses or were an intermediate on the pathway to large-genome evolution. However, the functional data led us to favor a model in which the relaxation of the interactions with the motif F arginine preceded the acquisition of the SDD in motif C; in the context of an ancestral 3Dpol-like enzyme, the E:SDD configuration is a slow and very high-fidelity polymerase that would likely reduce virus fitness and limit its emergence.

**Summary.** In this study, we addressed the molecular and mechanistic basis for the high speed and fidelity of the SARS-CoV-2 core replicase. Using a nsp8L7 fusion protein, we show that assembly of the (nsp12-nsp7/8-nsp8) core replicase complex is rate limited by the binding of the individual nsp8 molecule to the nsp12 polymerase fingers domain. The resulting complex is the fastest viral RdRP we have observed to date, with a processive elongation rate of 260 nt/s at 37° that is 3-fold faster than that of poliovirus polymerase. We have identified two noncanonical residues in the nsp12 active site that tailor the CoV enzyme to carry out large-genome replication: First, Ala547 in motif F replaces a canonical glutamate and allows for a doubling of replication rate by un tethering Arg555 in the active site, but this comes at a fidelity cost due to less restrictive NTP positioning during catalysis. Second, Ser759 in motif C restores fidelity by positioning the priming ribose for catalysis, in effect creating a new fidelity control point to compensate for the loss of NTP positioning. These structural and biochemical interactions suggest there were three major steps in the evolution of the nidovirus polymerase machinery (Fig. 7): (i) mutation away from the motif F glutamate to slightly increase elongation rates, (ii) the adoption of a SDD instead of GDD sequence in motif C as new fidelity control point via primer positioning, and (iii) further evolution of even faster elongation rates by adopting a motif F alanine that fully untethers the active-site arginine. The first two of
these events allow for stable maintenance of 13- to 16-kb arterivirus genomes, while the third enabled the rapid replication of larger 26- to 32-kb genomes but also required the acquisition of more powerful error-correcting ExoN functionality to maintain genomic stability. These findings further demonstrate ways in which the core structure and active-site closure mechanism shared by all positive-strand RNA virus polymerases can be evolutionarily tuned to optimize virus fitness.

MATERIALS AND METHODS

Protein expression and purification. A T7-based pET-26 plasmid with a N-terminal His6 tag followed by a TEV cleavage site was used to express codon optimized versions of the Wuhan-1 strain SARS-CoV-2 proteins. The nsp8L7 fusion proteins were made by adding a GSGSGS linker and nsp7 to the C terminus of nsp8. All proteins were expressed in E. coli BL21(DE3) CodonPlus cells (Agilent Technologies) and grown at 37°C to an optical density at 600 nm (OD600) of 0.8 prior to induction with 0.5 mM isopropyl-β-D-thiogalactopyranoside (IPTG). Nsp8, nsp7, and nsp8L7 and all N-terminal deletion constructs were expressed at room temperature for 18 h in NZCYM broth. Harvested cells were lysed at 18,000 lb/in² using a M-110L microfluidizer (Microfluidics Corp.) in a buffer containing 50 mM Tris, pH 8.0, 300 mM NaCl, 10 mM imidazole, 20% glycerol. The lysate was centrifuged at 17,000 rpm in a Beckman JA-20 rotor, and the supernatant was passed through 0.45- and 0.20-

RNA selection criteria. For the experiments described below, we have used different RNA designs to accommodate conflicting experimental constraints. For fluorescence polarization (FP)-based RNA-binding studies, we chose a short primer-template hairpin that provides both duplex and single-stranded RNA regions while being small enough to elicit a significant FP change upon protein binding. For the assembly and fidelity studies, we chose hairpin RNAs that would provide a stable primer-template duplex for initiation, yet were small enough to fully denature on gels so as to provide clear single-base resolution that was not complicated by internal base pairing that can lead to smearing. For the stopped-flow kinetic assays, we needed long-lived RdRP-RNA complexes that could be diluted to ~50 nM before being loaded into the instrument for 10 to 12 sequential sample injections (Fig. 4) over a 3- to 5-min time period without significant dissociation. For the picornaviral polymerases, we can preincorporate a few nucleotides to form stable stalled elongation complexes with very slow koff rates, but this does not work for CoV nsp12 (24), and we instead used a much longer RNA duplex section to gain stability via making full contact with the nsp8 sliding pole helices.

RNA binding. Fluorescence polarization (FP) RNA-binding assays were performed using a synthetic RNA hairpin with a 5′-GAAUGGUCUC-3′ single-stranded section labeled at its 5′ end with a fluorescein dye via a 6-carbon linker. Protein titrations were done with 20 nM RNA in binding buffer consisting of 50 mM HEPES (pH 7.0), 50 mM NaCl, 1 mM MgCl2, 4 mM TCEP, and 0.1% (vol/vol) NP-40 detergent. FP was measured in black 384-well polystyrene plates using a Perkin-Elmer Victor® multimode microplate reader. RNA dissociation constants were obtained by curve fitting the FP data to the quadratic form of the single-site binding isotherm.

Core replicase assembly. Assembly and initiation experiments were conducted by mixing final concentrations of 2.5 μM nsp12, 5 μM each nsp8 and nsp8L7, and 1 μM RNA in 50 mM HEPES, pH 7.0, 62 mM NaCl, 1 mM MgCl2, and 1 mM TCEP with various orders of addition and incubation times. The RNA for this assay is a synthetic (Integrated DNA Technologies, Coralville, IA) 8-base pair priming hairpin followed by various single-stranded template sequences as indicated in the figures. The RNA hairpin was labeled at its 5′ end with an IRdye 800RS NHS ester (Li-Cor Biosciences). The proteins were preincubated as indicated at 2× final concentrations and mixed with remaining components at t = 0, and time point samples were taken thereafter. Each time point is a 15-s elongation reaction with 50 μM NTPs (further diluting the replicase sample 2-fold) to assay for the amount of competent complex present at the specified assembly time. The reactions were manually quenched with 50 mM EDTA, 95% formamide, 0.02% bromophenol blue, and 0.02% xylene cyanol and analyzed by denaturing polyacrylamide gel electrophoresis using 7 M urea, 17% 19:1 acrylamide, 1× Tris-borate-EDTA (TBE) gels. Resolved RNA bands were imaged and quantified with a Li-Cor Odyssey CLx IR imager system and plotted as a ratio of elongated product species to the total amount of RNA.
Core replicase stability. The temporal stability of the nsp12-nsp8 complex was assessed with a large scale 300 µL assembly reaction of 20 µM nsp12 with 60 µM nsp8 and using gel filtration chromatography in 100 mM NaCl, pH 7, 10% glycerol to purify the complex as a 0.5-mL sample from the leading edge of the peak at ~12-mL retention volume. This purified complex was incubated at room temperature, and 50-µL aliquots were periodically reinjected onto the column over an 18-h time course (Fig. 3E). The areas of the overlapping nsp12-8 and nsp12 peaks in the resulting chromatograms were determined by mathematical peak fitting (PeakFit, Systat Software) and corrected for their different extinction coefficients (http://web.expasy.org/protparam) to calculate the relative concentrations present in the loaded sample aliquots.

Elongation kinetics. Stopped-flow fluorescence was used to measure kinetics of processive elongation on a three-piece annealed RNA with a 29-bp duplex primer and a 26-nt single-stranded template with a fluorescein placed at the 5′ end of the template. RNAs were synthesized by integrated DNA Technologies (Coralville, IA) and annealed by mixing all three strands and heating to 70°C followed by slow cooling at 1°C/min for 60 min in 66 mM KCl, 1 mM Tris, pH 8.0, and 0.1 mM EDTA. Replicase elongation complexes were generated by incubating 8 µM nsp12 with 10 µM nsp8 for 45 min at room temperature. 10 µM nsp8L7 was incubated with 6 µM RNA for 15 min prior to mixing 1:1 with the nsp12/8 complex and further incubated for 15 min. The replicase mix was then diluted 60-fold into 75 mM NaCl, 4 mM MgCl2, and 5 mM MES, pH 7.0, and 1 mM TCEP and immediately loaded into the Bio-Logic SFM-4000 titrating stopped-flow instrument with a MOS-500 spectrometer, and kinetics data were collected using equimolar mixtures of all four NTPs at various concentrations at 8°C (http://www.dnastars.com/). For the replicase assembly reaction of 20 µM nsp12 with 60 µM nsp8L7 with 2 µM RNA in 50 mM HEPES, pH 7.0, 50 mM NaCl, 1 mM MgCl2, and 1 mM TCEP that had been preincubated for 15 min. Replicase was assembled for an additional 15 min prior to the addition of 50 µM ATP and various concentrations of CTP for 15 s before manual quenching with 50 mM EDTA in 95% formamide, 0.02% bromophenol blue. For the experiments with poliovirus and coxsackievirus polymerases, 5 µM 3Dpol enzymes were preincubated with 1 µM RNA for 15 min prior to initiating burst elongation using the same buffers, nucleotide mixtures, and reaction times as for the assembled nsp12 complex. Note that we did not preinitiate to generate stalled 3Dpol elongation complexes for these experiments, i.e., the nsp12 and 3Dpol assays are done under identical conditions requiring first nucleotide addition followed by processive elongation and misincorporation. Monovalent salts can significantly affect RNA binding, and the salt concentrations listed above include accounting for any salt introduced with the proteins whose purification buffers contained 200 to 300 mM NaCl. Quenched elongation samples were resolved by denaturing polyacrylamide gel electrophoresis using 7 M urea, 17% 19:1 acrylamide, and 1× TBE and imaged with a Li-Cor Odyssey CLx IR imaging system. Misincorporation was quantified by integrating the RNA band intensities and calculating the ratio of misincorporation products to the total amount of elongated products, i.e., all cognate and noncognate incorporations, and plotting them against the CTP concentrations. The quantitation for CTP:A mismatch is based on the sum of the +5 and +9 products divided by the total RNA elongated, i.e., sum of the +3, +5, and +9 products, and for ATP:C, it is based on the +9 product divided by the sum of the +5 and +9 products, i.e., what fraction of the +5 is further elongated to yield +9. Both quantitation schemes analyze only the material that was elongated in the burst-phase assay.

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