The Nonstructural Protein 3 Protease/Helicase Requires an Intact Protease Domain to Unwind Duplex RNA Efficiently*

Received for publication, September 25, 2003, and in revised form, October 24, 2003
Published, JBC Papers in Press, October 29, 2003, DOI 10.1074/jbc.M310630200

David N. Frick‡§, Ryan S. Rypma‡, Angela M. I. Lam‡, and Baohua Gu¶
From the ‡Department of Biochemistry and Molecular Biology, New York Medical College, Valhalla, New York 10595 and the ¶Department of Biochemistry and Molecular Pharmacology, the Jefferson Center of Biomedical Research, Doylestown, Pennsylvania 18901

The nonstructural 3 (NS3) protein encoded by the hepatitis C virus possesses both an N-terminal serine protease activity and a C-terminal 3′–5′ helicase activity. This study examines the effects of the protease on the helicase by comparing the enzymatic properties of the full-length NS3 protein with truncated versions in which the protease is either deleted or replaced by a polystyridine (His tag) or a glutathione S-transferase fusion protein (GST tag). When the NS3 protein lacks the protease domain it unwinds RNA more slowly and does not unwind RNA in the presence of excess nucleic acid that acts as an enzyme trap. Some but not all of the RNA helicase activity can be restored by adding a His tag or GST tag to the N terminus of the truncated helicase, suggesting that the effects of the protease are both specific and nonspecific. Similar but smaller effects are also seen in DNA helicase and translocation assays. While translocating on RNA (or DNA) the full-length protein hydrolyzes ATP more slowly than the truncated protein, suggesting that the protease allows for more efficient ATP usage. Binding assays reveal that the full-length protein assembles on single-stranded DNA as a higher order oligomer than the truncated fragment, and the binding appears to be more cooperative. The data suggest that hepatitis C virus RNA helicase, and therefore viral replication, could be influenced by the rotations of the protease domain which likely occur during polyprotein processing.

The epidemic caused by infection by the hepatitis C virus (HCV)1 is still a global crisis despite recent therapeutic advancements (1). Because HCV cannot be conventionally cultivated in cell culture and the only host is the chimpanzee, the enzymes encoded by HCV have been studied intensely as targets for rational drug design. One key viral enzyme is the multifunctional nonstructural protein 3 (NS3), which possesses a serine protease activity and an ATPase function that fuels the ability of the protein to unwind RNA and DNA duplexes. Although it is clear that both the protease and helicase functions are necessary for viral replication (2), it is not clear whether the two functions, which reside in independent protein domains, cooperate in any manner (3).

To examine possible effects of the NS3 protease on its helicase function, the activities of the full-length NS3 protein were rigorously compared with the same protein lacking the protease and also with recombinant proteins in which the protease is replaced with other non-HCV peptides. The experiments were designed to uncover effects of the NS3 protease domain on its helicase function which are either specific or nonspecific. Nonspecific effects are defined as those that can be duplicated by peptides not derived from HCV NS3, whereas specific effects cannot.

All recombinant proteins used in this study (Fig. 1) were derived from the same infectious clone of HCV genotype 1a (4). NS3 spans amino acids 1027–1659 of the 3′–5′ amino acid long polyprotein encoded by HCV. At the N terminus resides the protease, which is made of two subdomains. The active site with its chymotrypsin-like catalytic triad is formed in a shallow cleft between the two protease subdomains. The NS3 protease is activated by the NS4A protein, which is translated immediately after NS3. The NS3-4A complex cleaves itself at the NS3-4A junction, and NS4A remains associative with NS3 (Fig. 1) as the complex cleaves the rest of the viral polyprotein. To do so, the protease would likely need to move away from the back of the helicase so that its active site is accessible to other peptides (5).

The helicase portion of NS3 is a three-domain Y-shaped molecule with two clefts separating the domains (6–8). The cleft between domains 1 and 2 is lined by several conserved motifs that are involved in ATP hydrolysis and are shared with other helicases. One strand of the nucleic acid substrate binds in the cleft that separates domain 3 from domains 1 and 2 (7). Fig. 1B shows the relative location of the protease when the C terminus of NS3 is bound in the protease active site (5). In this conformation, the protease active site and NS4A are far from the known RNA and ATP binding sites.

To analyze the helicase activities of the full-length protein, NS3 and NS4A were coexpressed as a single polypeptide that processes itself (9). The truncated helicase proteins lacking protease function (Fig. 1A) were based on those previously analyzed and include versions with N-terminal (10–12) and C-terminal (7, 13–16) polyhistidine tags (His tags) and a version in which the protease is replaced with a glutathione S-transferase (GST) protein (17). The presence of an intact protease domain dramatically enhances the ability of the protein to unwind RNA. However, the addition of an N-terminal His tag to the helicase fragment also enhanced RNA helicase activity, and the N-terminal GST-tagged protein unwind RNA almost as well as the full-length protein. Similar but smaller effects were seen in DNA unwinding assays and ssDNA trans-
Effects of NS3 Protease on HCV Helicase

**EXPERIMENTAL PROCEDURES**

**Materials**

RNase-free reagents were purchased from Ambion (Austin, TX), and nucleotides and nucleic acids were treated with the RNAse secure reagent (Ambion). Poly(U) RNA with an average length of 2,500 nucleotides was purchased from Sigma. DNA oligonucleotides were purchased from Integrated DNA Technologies (Coralville, IA), and their concentrations were determined from provided extinction coefficients.

**Protein Expression / Purification**

The recombinant proteins utilized in this study were all derived from the same HCV isolate (4) and possess the viral regions indicated in Fig. 1. The proteins are designated according to the type and location of their fusion partners. After purification, the following extinction coefficients, derived using the method described by Pace et al. (18), were used to determine protein concentrations: His-NS3-4A, 77.3 m"-1 cm"-1; His-Hel, 49.1 m"-1 cm"-1; His-Hel-His, 49.1 m"-1 cm"-1; GST-Hel-His, 87.4 m"-1 cm"-1; Cloning, expression, and purification of each are described below.

**His-NS3-4A**—PCR was used to amplify NS3-4A from the plasmid pCV-H77c, which was a generous gift from Dr. Jens Bukh (National Institutes of Health). The PCR primers added restriction sites to either end of the DNA and encoded the 5′-His tag depicted in Fig. 1. The sequence of the upstream primer was 5′-CAAGATCT ATG CAT CAC CAT CAC CAT GCC GGC GCC CCG ATC ACG GCC GAC GGC TAC GCC-3′ (BglII site underlined), and the sequence of the downstream primer was 5′-CAGAATTC TCA GCA CTC TTC CAT CTC ATC GAA C-3′ (EcoRI site underlined). The amplified DNA was digested with BglII and EcoRI and ligated into a similarly treated plasmid pVL1392 (Invitrogen). After verification of the appropriate DNA sequence, the resulting plasmid was recombined with the BacPAK system (Clontech) to generate a recombinant baculovirus. Baculovirus was plaque purified, amplified, and used for infection of SF9 cells for protein purification. The protein was isolated from SF9 cells using the protocol described by Sali et al. (9). Recombinant proteins were monitored during the purification procedure by Western blotting using antibodies raised against NS3 and NS4A.

**His-NS3**—To generate a recombinant HCV helicase fragment with both C-terminal and N-terminal His tags, DNA encoding the helicase was excised from plasmid p24Hel-1a (15), which encodes Hel-His, and inserted into the vector pET33a (Novagen, Madison, WI). Plasmid p24Hel-1a was digested with NheI and BamHI. The resulting smaller DNA fragment was purified and inserted using T4 DNA ligase into a similarly treated pET33a to generate the vector p33Hel-1a. For expression, p33Hel-1a was used to transform strain BL21(DE3), which contains T7 polymerase under the control of a lac promoter. The recombinant protein is under control of a T7(lac) promoter, which contains a lac operator between the T7 promoter and the transcription start site (19). After cells had grown to mid log phase in Luria-Bertani broth, helicase expression was induced with 1 mM isopropyl-β-D-thiogalactopyranoside. Cells were harvested and lysed by two passes through a French pressure cell, and the clarified soluble extract was fractionated as described previously for Hel-His (15).

**His-Hel**—To generate a HCV helicase protein with a single His tag on its N terminus, plasmid p33Hel-1a was modified using site-directed mutagenesis to introduce a stop codon at the end of the HCV NS3 "gene." The QuikChange site-directed mutagenesis kit (Stratagene) was used to modify p33Hel-1a using complementary mutagenic oligonucleotides: 5′-CCA CCT GCT AGC GCC GAA GGA GGA GGA GAC CCC ACC-3′ and 5′-CCG ATG CTC GTT ACC GCT GAG GCC GTG TCC GCT GAC-3′ (stop codon underlined). After the location of a stop codon was verified by DNA sequencing, the resulting plasmid (designated p33Helstop) was introduced into BL21(DE3) cells. Recombinant His-Hel was expressed and purified as described above for His-Hel-His.

**GST-Hel-His**—The GST-tagged HCV helicase was generated by excising the helicase encoding region from p24Hel-1a and inserting it into vector pET41a (Novagen). Plasmid p24Hel-1a was digested with the restriction enzymes NheI and BamHI. After the smaller of the two resulting DNA fragments was purified, it was ligated into pET41a that was digested with SpeI and BamHI (SpeI and NheI generate compatible cohesive ends), GST-Hel-His was then expressed in BL21(DE3) as described above. The GST-Hel-His protein was purified using the columns described for Hel-His (15) with the following modifications. After the nickel-nitriloacetic acid column, pooled fractions were loaded onto a 3-ml GST-bind™ column (Novagen). The column was washed with 50 ml of buffer A (50 mM HEPES, 1 mM EDTA, pH 7.5), and the protein was eluted with a gradient of the same buffer containing 0–20 mM reduced ATP binding site. In the complex, the C terminus of NS3 is bound in the active site of the protease, which is formed from a catalytic triad located assays. A more specific effect of the protease was seen when RNA unwinding was analyzed in the presence of a DNA trap, which sequesters excess helicase not bound to the RNA substrate. More efficient unwinding correlates with a different mode of interaction seen in direct DNA binding assays, in which twice as many protein monomers bind the same amount of ssDNA. Finally, ATPase assays show that the full-length helicase hydrolyzes ATP more slowly than the truncated proteins (in the presence of DNA or RNA), suggesting that the protease domain allows the helicase to fuel RNA unwinding more efficiently with ATP hydrolysis.
glutathione. Fractions containing HCV helicase were combined, precipitated with 60% ammonium sulfate, and purified further using gel filtration and DEAE chromatography as described previously for Hel-His (15).

**Unwinding Assays**

To generate substrates for helicase assays, two synthetic oligonucleotides were annealed by heating them to 95 °C and allowing them to cool slowly to room temperature. Before annealing, the shorter strand was 32P labeled using polynucleotide kinase. The duplex RNA substrate consisted of the RNA oligonucleotide 5'-GGCGCACCGACCCGGAG- C-3' annealed to H77c 3' UUUUUUUUUUUUUUUUURUU-3'. The DNA substrate consisted of a shorter strand DNA oligonucleotide 5'- 3P-GCCCTGCTGCGGTC- GCCA-3' annealed to a longer strand DNA oligonucleotide 5'-GGCGG- ACAGCAGCGAGG- TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT...
Fig. 2. Purified recombinant proteins. A, a 10% polyacrylamide gel containing 1% SDS with 1 μg each of the proteins used in this study: lane 2, His-NS3-4A, 5.786 kDa (NS4A), 68.350 kDa (His-N55); lane 3, His-Hel, 52.697 kDa; lane 4, His-Hel-His, 55.484 kDa; lane 5, Hel-His, 52.731 kDa; lane 6, GST-Hel-His, 79.682 kDa. B, to visualize the NS4A protein not seen in the 10% gel, 5 μg of the His-NS3-4A complex was separated on a 20% SDS gel and transferred to nitrocellulose membranes for Western analysis. Lane 1 shows a gel stained with Coomassie Blue, lane 2 shows a membrane probed with antibodies raised against NS4A, and lane 3 shows an identical membrane stained with antibodies raised against NS3.

Effects of NS3 Protease on HCV Helicase

The Protease Domain Aids RNA Unwinding—Previous studies using Hel-His have shown that it unwinds RNA poorly relative to DNA (15, 16), and when excess DNA is added upon initiation to reactions containing Hel-His, virtually no RNA unwinding is detected (15). Although a preference for DNA has been long established for HCV helicase (32), the absence of RNA helicase activity in the presence of an enzyme trap (such as excess DNA) starkly contrasted results published previously by other authors (25, 31). The ability of a helicase to unwind DNA (or RNA) in the presence of a trap is indicative of a processive reaction because if enzyme molecules dissociate after only a few base pairs are unwound then no reaction will be detected. The trap prevents enzymes not initially bound to substrate from participating in the reaction (33). Thus, our previous results suggest that Hel-His is a processive DNA helicase but not a processive RNA helicase (15, 16), which is a very different conclusion than that of Pang et al. (25), who showed the HCV helicase is processive on both DNA and RNA. The study by Pang et al. (25) differed from ours (16) in two ways. First, the helicases were isolated from different HCV strains, and second, Pang et al. (25) used full-length NS3 proteins, whereas we used the truncated protein Hel-His (16). This study was therefore initiated using proteins with no genotypic variation to test whether the protease in fact aids RNA unwinding. The ability of four of the proteins studied here to unwind duplex RNA is shown in Fig. 3. RNA unwinding by Hel-His was analyzed previously (15). The RNA helicase assays in Fig. 3 are done in the absence of a trap but under conditions in which no RNA reannealing occurs. The duplex RNA substrate consists of a 5′-32P-labeled 18-nucleotide-long RNA oligonucleotide annealed to a 38-nucleotide-long oligonucleotide in such a manner to create a substrate with a 3′-ssRNA tail. This tail is required for unwinding. None of the helicases studied here unwinds substrates with either blunt ends or with only a 5′-ssRNA tail. In these assays, the RNA and helicase are combined and incubated for 30 min to allow efficient substrate loading. Reactions are initiated with ATP, quenched at various times, and analyzed on nondenaturing polyacrylamide gels.

All proteins with peptides attached to the N terminus of domain 1 of HCV helicase unwind RNA better than Hel-His, which lacks an N-terminal attachment (Fig. 3). When the initial rates of RNA unwinding by the N-terminally tagged enzymes (Fig. 3, A-D) are compared with rates derived from the same experiments conducted with Hel-His, dramatic differences are apparent (Fig. 3E). Less of these proteins are required to initiate unwinding, and RNA is unwound faster and almost to completion. The initial rates of RNA unwinding can be analyzed to yield a maximum unwinding rate (Vmax) and a concentration of helicase which leads to 50% maximum activity. Estimates of these parameters can be obtained by fitting the data in Fig. 3E to the Michaelis-Menten equation. However, because the data better describe sigmoid curves, such an analysis would be misleading. The data were therefore fit to a sigmoidal dose-response curve with a variable slope. The top of these curves provides an estimate of Vmax and the midpoint provides an EC50 value. Such an analysis (Table I) reveals that His-NS3-4A unwinds RNA most efficiently with an EC50 of 54 nM and a Vmax of 2.6 nM/min. GST-Hel-His unwinds RNA at a slightly faster maximum rate but with a higher EC50. This suggests that the protease is structurally important for RNA unwinding, but it can be essentially replaced by a protein of entirely different origin. Surprisingly, even the proteins in which the protease is replaced with a short His tag unwind RNA almost as well as the full-length NS3 protein (Table I).

Unlike Hel-His, the other proteins analyzed here unwind RNA in the presence of a DNA trap (Fig. 4). In these assays, reactions were initiated with ATP and various amounts of a DNA oligonucleotide that consists of the same sequence as the shorter strand in the RNA duplex (except with Ts replacing Us). The trap binds both free enzyme not bound to substrate and enzyme that dissociates from the substrate before the
reaction is terminated. In other words, only a single reaction cycle of enzyme and substrate should be theoretically observed in the presence of trap DNA. In the presence of 20 times more trap than enzyme, all of the helicases except Hel-His could unwind a significant amount of RNA. The initial rates of RNA unwinding in reactions catalyzed by His-NS3-4A (33), His-Hel (33), His-Hel-His (33), and GST-Hel-His (33), are compared with the data obtained with Hel-His (data are from Ref. 16 (C)). Data are fit to a sigmoidal dose-response curve with variable slope using nonlinear regression (using Graphpad Prism, version 4.0). The resulting maximum velocities and EC50 values are listed in Table I.

TABLE I

<table>
<thead>
<tr>
<th>Protein</th>
<th>RNA (37 °C)</th>
<th>DNA (23 °C)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Vmax</td>
<td>EC50</td>
</tr>
<tr>
<td></td>
<td>nM/min</td>
<td>nM</td>
</tr>
<tr>
<td>His-NS3-4A</td>
<td>2.6 ± 0.2</td>
<td>54 ± 5</td>
</tr>
<tr>
<td>His-Hel</td>
<td>2.4 ± 0.1</td>
<td>95 ± 3</td>
</tr>
<tr>
<td>His-Hel-His</td>
<td>1.8 ± 0.1</td>
<td>53 ± 2</td>
</tr>
<tr>
<td>Hel-His</td>
<td>0.8 ± 0.1</td>
<td>182 ± 5</td>
</tr>
<tr>
<td>GST-Hel-His</td>
<td>2.9 ± 0.1</td>
<td>74 ± 5</td>
</tr>
</tbody>
</table>

To compare properly the efficiency with which each protein unwinds DNA or RNA, DNA unwinding assays were repeated at 23 °C (Fig. 5B). Each of the proteins clearly prefers DNA except His-NS3-4A, again highlighting a specific role for the protease in RNA unwinding (Table I).

We have shown previously that DNA unwinding catalyzed by Hel-His is not particularly sensitive to the presence of a DNA prefered to DNA unwinding.
trap (16). Not surprisingly, DNA unwinding catalyzed by the other proteins is likewise insensitive to a trap (data not shown), suggesting that they are all processive DNA helicases.

**DNA Translocation Assays**—Morris et al. (21) have shown that in addition to unwinding RNA and DNA, the full-length HCV genotype 1b NS3 protein is able to strip streptavidin from biotin-labeled DNA. Because the HCV helicase can only strip streptavidin from the 5’-end of DNA, these assays provide evidence that the protein translocates only in a 3’–5’ direction. We recently confirmed these experiments using Hel-His, but because our studies used a protein derived from genotype 1a, no firm conclusions were reached regarding the role of the protease in DNA translocation (16). We have therefore repeated these studies using all the proteins described here (Fig. 6). The new data show that the protease domain accelerates the translocation rate of the protein on ssDNA but that this effect can be substituted entirely by a GST fusion peptide (GST-Hel-His). An N-terminal His tag does not appear to aid translocation because His-Hel moves only somewhat faster than Hel-His, whereas His-Hel-His moves slightly slower. The initial translocation rates of each protein seem to obey Michaelis-Menten kinetics (Fig. 6B), unlike initial rates of RNA unwinding (Fig. 3E), or DNA unwinding (Fig. 5A). This may simply be because the enzyme and substrate are in a state of rapid equilibrium or because the reactions are not processive. The addition of a DNA trap to these reactions abolishes any detectable dissociation of ssDNA from streptavidin (data not shown).

**Binding Assays**—The issue of whether or not helicases unwind nucleic acid duplexes as monomers, dimers, or higher order oligomers is still hotly debated. Numerous studies have concluded that HCV helicase functions as a monomer (10, 34, 35), whereas many others have presented clear evidence for oligomerization (17, 36, 37). It has been proposed that these differences might be because the protease region is required for stable oligomerization (17, 37), but this has not yet been directly demonstrated. Although all of the proteins here migrate mainly as monomers on both gel filtration columns and native polyacrylamide gels, the sigmoidal dependence of unwinding on protein concentration suggests that interprotein interactions might be critical for RNA unwinding (Fig. 3E). To address this...
issue further, binding assays were performed to quantify the interaction of each helicase with ssDNA. Previously, intrinsic protein fluorescence has been used to examine ssDNA interaction with HCV helicase (15, 16, 22, 35, 38). All of these studies used recombinant proteins lacking the protease domain and exploit the fact that ssDNA binds near a solvent exposed tryptophan to quench intrinsic protein fluorescence (7). Previously, we have performed these studies using Hel-His (see Fig. 8 of Ref. 16). However, it was extremely difficult to repeat these experiments with the other proteins examined here because the intrinsic protein fluorescence of each changed more rapidly in the absence of added oligonucleotide. As shown in Fig. 7A the intrinsic protein fluorescence of Hel-His decreases only about 5% over 10 min. Others have reported similar drifts (35, 38), which were corrected for by performing “blank” titrations where buffer was added at the same intervals as the additions of ssDNA. The data were then corrected for this minor drift.

This drifting was more obvious with proteins containing peptides attached to the N terminus of the helicase (Fig. 7A). The intrinsic protein fluorescence of the full-length protein, His-NS3-4A, and the N-terminal His-tagged proteins decreased almost 20%, whereas that of GST-Hel-His decreased almost 40%. To investigate the physical basis for this fluorescence drift, the experimental conditions were varied systematically. The drift was independent of pH, salt concentration, buffer composition, temperature, or divalent metal ion concentration. Only the concentration of Tween 20 affected drift rates, with faster changes in intrinsic protein fluorescence occurring at lower concentrations of detergent. The data in Fig. 7 were obtaining in the presence of 0.2% Tween 20. Higher detergent concentrations were not used because background fluorescence from higher concentrations of Tween 20 made observations of intrinsic protein fluorescence difficult.

There are two possible explanations for the drift in intrinsic protein fluorescence: either the protein is slowly changing the conformation (denaturing) or aggregating. Two lines of evidence support the idea that the changes in intrinsic protein fluorescence reflect protein oligomerization rather than denaturation. First, no changes in specific activity in either helicase or ATPase assays were measured when any of the helicase proteins were incubated for several hours at temperatures as high as 37 °C. Second, oligomerization of the HCV helicase has...
been observed by others using cross-linking (35), yeast two-hybrid assays (17, 39), and gel filtration chromatography (9, 17). Although the vast majority of the proteins studied here migrate as monomers (or in the case of His-NS3-3A, a heterodimer), a small amount of the HCV proteins can be detected in the void volume of the columns, as was first reported by Sali et al. (see Fig. 3 of Ref. 9).

The previously reported oligomerization of HCV helicase has been reported to be greatly enhanced by the presence of oligonucleotides. The presence of ssDNA oligonucleotides allows for more apparent dimers, trimers, tetramers, and pentamers to be detected by cross-linking (36) and gel filtration chromatography (17). Thus, if the changes in protein fluorescence reflect a slow oligomerization, then the rates of fluorescence change should be accelerated in the presence of a ssDNA oligonucleotide. To test this hypothesis, the experiments in Fig. 7A were repeated in the presence of DNA. Indeed, added oligonucleotides accelerated the rate of intrinsic protein fluorescence decrease for all proteins. The data obtained with GST-Hel-His are shown in Fig. 7B. When ssDNA is present, it not only quenches fluorescence, but it also accelerates drift. The intrinsic protein fluorescence stabilizes earlier in the presence of DNA than in the absence of DNA (Fig. 7B). Although this observation supports the idea that fluorescence changes reflect oligomerization, it greatly complicates the analysis of DNA binding to HCV helicase when intrinsic protein fluorescence is monitored.

Because quenching of intrinsic protein fluorescence provided a poor method to examine helicase DNA interactions (for proteins other than Hel-His) several fluorescently labeled DNA molecules were screened to find one that displayed a large fluorescence change upon protein binding. We found that the molecule that gave the best signal was an 18-nucleotide-long oligonucleotide with a fluorescein moiety attached to its 3′-end (F18). To show that the fluorescence itself did not influence the interaction between helicase and the oligonucleotide, its binding to Hel-His was analyzed by monitoring changes of intrinsic protein fluorescence (Fig. 7C). F18 and the same oligonucleotide lacking the fluorescein (18-mer) bound similarly both in the presence and absence of the nonhydrolyzable ATP analog ADP(BeF3). As was seen previously, DNA binds HCV helicase about 10-fold weaker in the presence of ADP(BeF3) (16, 22).

This has been interpreted to mean that ATP binding allows the protein to slide along DNA (22). Using this assay, the apparent $K_D$ for the 18-mer in the absence of ADP(BeF3) is $1.9 \pm 0.3$ nM, whereas the $K_D$ for F18 is $2.5 \pm 0.3$. These values are not significantly different. Likewise, the $K_D$ for the 18-mer with ADP(BeF3) is $26 \pm 2$, and the $K_D$ for F18 is similar at $30 \pm 2$.

The shapes of the binding isotherms in Fig. 7C clearly indicate tight binding, but interestingly, the inflection points do not match the total amount of protein monomers in solution (50 nM, as measured from the protein extinction coefficient). To fit a standard equation describing the binding of a ligand to a protein, the ligand concentration must be adjusted by a factor, $n$ (see Equation 1, which was derived from Equations 2, 3, and 4 of Ref. 35). The factor, $n$, can be interpreted as the number of protein monomers that bind a single oligonucleotide. Although a value of $n = 3.7$ was determined from the data in Fig. 7C by nonlinear regression analysis, a similar value can be determined qualitatively simply by examining the graph. All curves inflect around 13.5 nM DNA, which is about the amount of DNA necessary to bind all 50 nM protein. Hence, three or four proteins monomers bind a single template, or they each cover about 5 nucleotides of the 18-mer.

Changes in F18 fluorescence upon addition of all the helicases studied here except GST-Hel-His is shown in Fig. 8A. In control experiments, neither addition of buffer nor a Hel-His mutant (F444A) lacking detectable ATPase, helicase, or binding activity (16) produced any changes in F18 fluorescence. GST-Hel-His did not yield analyzable data in this assay because after fluorescence increased upon protein addition, fluorescence began to decrease slowly. Fluorescence of the F18 complex made with the other proteins was much more stable. The data obtained with Hel-His at multiple oligonucleotide concentrations (Fig. 8D) fit an equation similar to the one used to analyze changes in intrinsic protein fluorescence to yield.

![Fig. 8. Analysis of ssDNA binding using a fluorescein-labeled oligonucleotide.](image)

The fluorescence of F18 was monitored at an excitation wavelength of 492 nm and an emission of 518 nm in the absence and the presence of various concentrations of various HCV helicase constructs. A change in fluorescence ($\Delta F$) when 2 nM F18 is titrated with His-NS3-4A ( ), His-Hel ( ), His-Hel-His ( ), or Hel-His ( ). The other three panels show how raw fluorescence of 0.5 nM ( ), 0.75 nM ( ), 1 nM ( ), 1.5 nM ( ) or 2 nM ( ) F18 changes upon addition of His-NS3-4A ( ), His-Hel ( ), or Hel-His ( ). Data are fit to Equation 2 using global nonlinear least squares analysis to yield the values listed in Table II.
ATP analog ADP(BeF₃) was fit to Equation 2 to yield an apparent dissociation constant and a factor D

values for $K_D$ and $n$ (Table II). However, when the same experiments are done with His-NS3-4A (Fig. 8B), His-Hel (Fig. 8C) or His-Hel-His (not shown), the resulting isotherms look very different from those obtained with Hel-His (Fig. 8D). Binding isotherms obtained with His-Hel-His (not shown) were virtually identical to those obtained with His-Hel. The isotherms obtained with His-Hel and His-Hel-His are S-shaped at low protein concentrations, very steep, and have clearly defined inflection points. The inflection points of His-Hel are at twice the protein concentration seen with Hel-His (compare Fig. 8, C and D), indicating that twice the amount of protein is required to saturate the binding sites on the DNA. Isotherms obtained with His-NS3-4A inflect in the same region as those obtained with His-Hel (Fig. 8C). The data are fit to the simple binding model described in Equation 2 to yield the parameters described in Table II.

Table II also summarizes the results of the same set of experiments when they were conducted in the presence of the nonhydrolyzable ATP analog ADP(BeF₃). Importantly, both Hel-His and His-NS3-4A appear to release ssDNA upon ATP binding, indicating that the full-length enzyme could function as a helicase are fueled by the hydrolysis of ATP. In all the helicase assays described above, no reactions were detected in the absence of ATP. Because the various proteins studied here unwind at different rates, it is conceivable that these differences are caused by different rates of ATP hydrolysis. Because HCV helicase hydrolyzes ATP in the absence of nucleic acids, and the ATPase rate is stimulated by DNA (and RNA), five different kinetic constants can be defined: the $K_m$ and $V_{max}$ in the presence and absence of nucleic acid, and the concentration of nucleic acid that yields 50% $V_{max}$ ($K_{DNA}$). These factors were determined by titrating each helicase with ATP in the absence (Fig. 9A) and presence (Fig. 9B) of saturating amounts of poly(U) RNA (the polymer that most efficiently stimulates ATP hydrolysis (40)), and by measuring ATP hydrolysis at saturating ATP concentrations at different concentrations of a DNA oligonucleotide (Fig. 9C). The resulting constants are summarized in Table III.

Because ATP hydrolysis fuels helicase action, one might reasonably assume that those proteins that efficiently unwind RNA also hydrolyze ATP the most rapidly. Surprisingly, this is

<table>
<thead>
<tr>
<th>Protein</th>
<th>$K_D$ (nM)</th>
<th>$n$ protein/DNA</th>
<th>$K_D$ (ADP(BeF₃))</th>
<th>$n$ protein/DNA</th>
</tr>
</thead>
<tbody>
<tr>
<td>His-NS3-4A</td>
<td>0.059 ± 0.029</td>
<td>6.7 ± 0.3</td>
<td>3.4 ± 0.5</td>
<td>6.6 ± 1.0</td>
</tr>
<tr>
<td>His-Hel</td>
<td>0.16 ± 0.06</td>
<td>7.0 ± 0.5</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>His-Hel-His</td>
<td>0.11 ± 0.08</td>
<td>8.2 ± 0.8</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Hel-His</td>
<td>0.23 ± 0.3</td>
<td>2.9 ± 0.2</td>
<td>3.8 ± 0.25</td>
<td>3.0 ± 0.1</td>
</tr>
</tbody>
</table>

* ND, not determined.

**TABLE II** ssDNA binding

Effects of NS3 Protease on HCV Helicase

Fluorescence data obtained when oligonucleotide F18 (Fig. 8) is titrated with HCV helicase in the presence and absence of the nonhydrolysable ATP analog ADP(BeF₃) is fit to Equation 2 to yield an apparent dissociation constant and a factor $n$, which describes the number of helicase monomers bound to a single oligonucleotide.

**FIG. 9.** ATP hydrolysis. Initial rates of ATP hydrolysis were measured at various ATP concentrations in the absence (A) and presence of 1 mg/ml poly(U) RNA (B). Rates (nM/s) were divided by the total amount of enzyme in the assay (nM) to compare the activities of His-NS3-4A (□), His-Hel (○), His-Hel-His (△), Hel-His (●), and GST-Hel-His (×). Data are fit to the Michaelis-Menten equation to yield the parameters summarized in Table III. C. Specific activity of ATP hydrolysis was measured at 4 mM ATP at various concentrations of a 20-nucleotide homopolymer of deoxyxymidine. Data are fit to Equation 3 to determine $K_{DNA}$, which is reported in Table III.
Effects of NS3 Protease on HCV Helicase

### DISCUSSION

Until now it has been generally assumed that the HCV NS3 protease and helicase are two independent units that are simply combined into a single multifunctional protein. Most previous studies have not noted differences in helicase (11, 23, 41), RNA binding (42), or RNA replication assays (24) when NS3 was truncated to delete the protease. In early studies Heilek and Peterson (11), concluded that the protease domain had little or no effect on helicase activity. In fact, in their assays, the full-length protein seemed to unwind RNA more slowly. Likewise, Gallinari et al. (23) concluded that there were no significant differences between the full-length protein and the isolated helicase domain in either ATPase or RNA unwinding assays. Both full-length NS3 and a truncated helicase fragment have been shown specifically to bind sequences in the HCV genome, and the protease is apparently not required for this interaction (42). In another interesting study that showed an ability of NS3 to stimulate RNA synthesis catalyzed by the HCV NS5B RNA-dependent RNA polymerase, both full-length and truncated NS3 behaved similarly (24). In contrast to these studies, Howe et al. (43) reported that a single chain recombinant protein where NS4A is attached to the N terminus of NS3 was more active in helicase assays than either NS3 or a truncated protease. That single chain NS4-NS3 protein was later crystallized and its structure is shown in Fig. 1B. When a molecular model of the single chain NS4-NS3 protein (5) is compared with one of a helicase fragment with a bound DNA oligonucleotide (7), it is clear that the protease active site and its NS4A cofactor are each more than 30 Å from the known DNA binding site and putative ATP binding site (Fig. 1B). The protease and helicase are connected via a flexible linker, and only a few residues in the protease domain contact residues in the helicase domain.

Although no role for the protease in helicase action has been identified previously, recently the NS4A protein has been suggested to help load helicase on RNA (44). This model is based mainly on the report by Pang et al. (25) that the presence of NS4A reduces the amount of preincubation time required for maximal rates of unwinding in the presence of a DNA trap. Also supporting this idea is the observation that when NS4A is covalently tethered to NS3, the helicase activity appears to be enhanced (43). It is therefore possible that the specific effects seen here are not caused by the protease, but instead NS4A. However, we are skeptical that the effects seen here are the result of NS4A for a couple of reasons. First, we did not find that the preincubation time influenced the RNA (or DNA) helicase activity of any of the proteins studied here (data not shown). Second, all assays were performed at low ionic strength, where NS4A might be dissociated from NS3. NS3 and NS4A bind via hydrophobic interaction stabilized by the presence of salt. When salt is absent, NS4A no longer stimulates HCV protease, indicating that it may be dissociated (45). Further examination of the role of NS4A will require purified NS4A which can be quantitatively added back to helicase assays. Unfortunately, this will be a very difficult experiment because of the insoluble nature of the NS4A protein.

The experiments described here clearly delineated effects of the NS3 protease domain on its helicase function which were discussed as either specific or nonspecific effects. For example, an NS3 protein lacking the protease domain unwinds duplex RNA slower than the full-length protein, but this rate can be accelerated by either the presence of an N-terminal His tag or GST tag (a nonspecific effect). On the other hand, no fusion protein can enable the helicase fragment to unwind RNA efficiently in the presence of a DNA trap (a specific effect). Of course, it is recognized these supposedly specific effects might

---

### TABLE III

**Kinetic analysis of ATP hydrolysis**

<table>
<thead>
<tr>
<th>Protein</th>
<th>Basal</th>
<th>RNA stimulated</th>
<th>K&lt;sub&gt;DNA&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>V&lt;sub&gt;max&lt;/sub&gt;/&lt;E&lt;sub&gt;T&lt;/E&gt;</td>
<td>K&lt;sub&gt;m&lt;/sub&gt;</td>
<td>V&lt;sub&gt;max&lt;/sub&gt;/&lt;E&lt;sub&gt;T&lt;/E&gt;</td>
</tr>
<tr>
<td></td>
<td>s&lt;sup&gt;-1&lt;/sup&gt;</td>
<td>µM</td>
<td>s&lt;sup&gt;-1&lt;/sup&gt;</td>
</tr>
<tr>
<td>His-NS3-4A</td>
<td>1.5 ± 0.2</td>
<td>230 ± 72</td>
<td>15 ± 1.3</td>
</tr>
<tr>
<td>His-Hel</td>
<td>3.0 ± 0.1</td>
<td>2.8 ± 0.37</td>
<td>56 ± 3.0</td>
</tr>
<tr>
<td>His-Hel-His</td>
<td>2.4 ± 0.2</td>
<td>11 ± 2.4</td>
<td>33 ± 1.2</td>
</tr>
<tr>
<td>Hel-His</td>
<td>1.8 ± 0.1</td>
<td>21 ± 0.42</td>
<td>86 ± 3.0</td>
</tr>
<tr>
<td>GST-Hel-His</td>
<td>0.56 ± 0.05</td>
<td>61 ± 15</td>
<td>43 ± 1.4</td>
</tr>
</tbody>
</table>

Initial rates of ATP hydrolysis (Fig. 9) in the presence and absence of 1 mg/ml poly(U) RNA were fit to ATP concentration using the Michaelis-Menten equation to yield V<sub>max</sub> and K<sub>m</sub> values. Also shown is the K<sub>DNA</sub>, defined as the DNA concentration needed to reach half of the V<sub>max</sub> seen at saturating DNA levels.
in fact be nonspecific because non-HCV peptides that were not tested might duplicate these effects.

One can speculate about how the protease-NS4A complex might nonspecifically influence the helicase based on the molecular model shown in Fig. 1B. All helicase fragments used here begin with a loop that extends along the back of the helicase connecting to domain 1 (red in Fig. 1). It is possible that this region that is not resolved in the structures of the helicase fragments (6–8), when not anchored to the back of the helicase, allows the protein to assume conformations that do not bind RNA tightly during helicase movements. The protease or fusion peptides might simply act like an anchor. Specific effects of the protease-NS4A complex are more difficult to explain. Perhaps a groove between the protease and helicase domain 2 provides a binding site for the strand of RNA not yet seen in the published molecular models. We plan to test some of these ideas using structure-based site-directed mutagenesis.

The results presented here also explain why many prior studies have only noted monomers of the HCV helicase, whereas others have detected dimers or higher order oligomers. As shown in Fig. 8 and Table II, the numbers of helicase monomers binding to a single oligonucleotide depend on the monomers binding to a single oligonucleotide depend on the monomers binding to a single oligonucleotide. It is also possible that inhibitors of the protease could lock the protein in a conformation that unwinds RNA poorly and therefore inhibit viral replication by both inhibiting polyprotein processing and RNA replication.

Clearly, the full-length and truncated versions unwind RNA very differently, and the activity of the isolated helicase fragments can be greatly influenced by the nature of the fusion protein tethered to its N terminus. This means that many of the general conclusions of numerous studies that utilized recombiant proteins encoding only fragments of the NS3 protein will need to be reconsidered, and many of the more rigorous studies will need to be repeated with the full-length NS3 protein.

Acknowledgments—We thank Fred Jaffe and Ruth Gallagher for valuable technical assistance.

REFERENCES