Motif V regulates energy transduction between the flavivirus NS3 ATPase and RNA-binding cleft

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The unwinding of dsRNA intermediates is critical for the replication of flavivirus RNA genomes. This activity is provided by the C-terminal helicase domain of viral nonstructural protein 3 (NS3). As a member of the superfamily 2 (SF2) helicases, NS3 requires the binding and hydrolysis of ATP/NTP to translocate along and unwind double-stranded nucleic acids. However, the mechanism of energy transduction between the ATP- and RNA-binding pockets is not well-understood. Previous molecular dynamics simulations conducted by our group have identified Motif V as a potential "communication hub" for this energy transduction pathway. To investigate the role of Motif V in this process, here we combined molecular dynamics, biochemistry, and virology approaches. We tested Motif V mutations in both the replicon and recombinant protein systems to investigate viral genome replication, RNA-binding affinity, ATP hydrolysis activity, and helicase-mediated unwinding activity. We found that the T407A and S411A substitutions in NS3 reduce viral replication and increase the helicase-unwinding turnover rates by 1.7- and 3.5-fold, respectively, suggesting that flaviviruses may use suboptimal NS3 helicase activity for optimal genome replication. Additionally, we used simulations of each mutant to probe structural changes within NS3 caused by each mutation. These simulations indicate that Motif V controls communication between the ATP-binding pocket and the helical gate. These results help define the linkage between ATP hydrolysis and helicase activities within NS3 and provide insight into the biophysical mechanisms for ATPase-driven NS3 helicase function.

Arthropod-borne flaviviruses, such as yellow fever virus, Japanese encephalitis virus, Zika virus, West Nile virus (WNV), and dengue virus, are a major health concern in the tropical and subtropical regions of the world (1, 2). Infection from these viruses causes disease symptoms ranging from flu-like illness to encephalitis, hemorrhagic fever, coma, and potentially death (3). Over half of the world population is at risk for infection from one or more of these viruses (4, 5). Dengue virus, specifically, infects around 50 million people each year, and of those individuals, 20,000 contract dengue hemorrhagic fever leading to their mortality (6). Additionally, WNV over the past 20 years within the 48 continental United States has around 50,000 clinical infections with a 5% mortality rate (7, 8) (and see https://www.cdc.gov/westnile/statsmaps/index.html (accessed September 12, 2019)). Currently, there are no approved antiviral drugs for treating flaviviral infections, and the vaccines in circulation, like the yellow fever vaccine, are not readily available worldwide for most flaviviruses (10–12).

To develop new antiviral treatments (drugs and vaccines) for these viruses, a fundamental understanding of how these flaviviruses replicate is required.

Flaviviruses (Flaviviridae family) have a positive-sense single-stranded RNA (ssRNA) genome that is ~11 kb in length. The viral RNA genome is translated into a single polyprotein that is subsequently cleaved by host and viral proteases into three structural proteins (C, prM, and E) and eight nonstructural proteins (NS1, NS2A, NS2B, NS3, NS4A, 2K, NS4B, and NS5) (13, 14). The structural proteins contribute to the formation of mature viral particles, whereas the nonstructural proteins are responsible for replication of the viral genome and protecting the virus from attacks by the host cell innate immune system (13). Once the viral proteins are post-translationally processed, the positive-sense genomic ssRNA is used as a template to create a negative-sense anti-genomic RNA, forming a dsRNA intermediate complex (15, 16). The nascent negative-sense ssRNA serves as a template strand for producing new positive-strand RNAs that can be packaged into viral particles, translated into new proteins, or interfere with the RNA decay pathway (17, 18). Therefore, the unwinding of dsRNA intermediate to produce a free negative ssRNA for positive-strand synthesis is a critical component for the replication of flavivirus RNA genomes (15). This function is achieved by the C-terminal helicase domain of NS3.

Helicases are ubiquitous enzymes that are classified into superfamilies based on their primary structure and highly conserved motifs (19–21). NS3 helicase (NS3h) is a member of superfamily 2 (SF2) helicases (22). The structure of NS3h consists of three subdomains; subdomain 1 and subdomain 2 are...
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Figure 1. Eight structural motifs of Flavivirus NS3 helicases are highly conserved. The structural motifs (Motifs I, Ia, II, III, IV, IVa, V, and VI) are located between the ATP-binding pocket and RNA-binding cleft. Each motif is highlighted as follows: Motif I in dark green; Motif Ia in blue; Motif II in orange; Motif III in royal blue; Motif IV in purple; Motif IVa in magenta; Motif V in lime green; and Motif VI in red.

RecA-like structures that are highly conserved across SF2 helicases (23). The third subdomain is unique to the viral/DEAH-like subfamily of SF2 helicases and interacts with subdomains 1 and 2 to form the RNA-binding cleft. NS3h is a multifunctional enzyme, possessing three enzymatic activities: RNA helicase, nucleoside triphosphatase (NTPase), and RNA 5' triphosphatase (RTPase) (24–29). Previous studies have shown that the two latter activities share a catalytic active site where NTP binds between subdomain 1 and subdomain 2 (26, 30–32). The RNA helicase active site is distinct from that of the NTPase/RTPase active site, located at the helical gate between subdomain 2 and subdomain 3 of NS3 (33–35). The helical gate and β-wedge are responsible for splitting the dsRNA intermediate into two ssRNA strands: the positive-sense viral ssRNA and the negative-sense template ssRNA (33). Single-molecule and structural biology studies have suggested that the negative-sense ssRNA template enters the RNA-binding cleft, where the helicase utilizes the energy produced from the hydrolysis of one ATP molecule to power translocation and unwinding of the dsRNA intermediate one base at a time (15, 16, 32, 36–40). A fundamental unanswered question within the SF2 helicase field is how ATP hydrolysis causes structural changes within the helicase to result in translocation and unwinding of dsRNA intermediates.

Between the ATPase active site and the RNA-binding cleft, there are eight structural motifs (Motifs I, Ia, II, III, IV, IVa, V, and VI) that are highly conserved in the viral/DEAH-like subfamily of SF2 helicases (Fig. 1) (22). These structural motifs play a critical role in substrate binding and are responsible for enzymatic activities within the helicase. One of these motifs, Motif V, was reported by Davidson et al. (24) as a potential link between the ATP-binding pocket and the RNA-binding cleft due to strong correlations between residues within Motif V and both binding pockets found within the helicase. Additionally, Mastrangelo et al. (34) reported that Motif V may be one of the main components of the driving force pulling the helicase along the ssRNA due to conformational changes initiated by the binding of ATP, which propagated through Motif V opening the helical gate. At least two residues within Motif V have direct interactions with either the ATP hydrolysis active site or the bound ssRNA molecule. Gly414 coordinates with the lytic water found within the ATPase active site, and Thr408 interacts with the phosphate backbone of the bound ssRNA (24). In addition to these interactions, Motif V has strong coupling between the other highly conserved motifs that interact with the ATP-binding pocket and the RNA-binding cleft, suggesting that Motif V may be central to the communication between these two binding pockets (24, 41). Therefore, we investigated the role of Motif V as a critical link between the two binding pockets. We utilized a combination of all-atom molecular dynamics simulations, biochemical assays, and virological assays to understand the role of Motif V in internal enzymatic communication leading to helicase function. Our results show that residues within Motif V control not only the enzymatic activities of the helicase but also the ability of the flaviviruses to replicate in vitro. This is due to the importance of the secondary structure of Motif V, as indicated through molecular dynamics simulation.

Results

Motif V mutants affect viral genome replication

As discussed previously by Davidson et al. (24), Motif V may play a role in the communication between the ATP-binding pocket and the RNA-binding cleft/helical gate. As one of the highly conserved motifs in SF2 helicases, Motif V consists of 13 residues spanning position 404 through position 416 (Fig. 2A). Eight of these residues are 100% conserved across all flaviviruses, both mosquito and tick-borne (Fig. 2B). The other five residues are highly conserved in flaviviruses that infect insects and mammals and are more variable in tick-borne restricted flaviviruses (Table S1). Interestingly, the conserved residues across all of the flaviviruses interact with either the ssRNA or ATP substrates. These highly conserved residues include Thr408, Asp409, Ile410, Glu412, Met413, Gly414, Ala415, Asn416. Of the mostly conserved residues, Thr407 and Ser411 mainly interact with each other through a hydrogen bond that is hypothesized to stabilize the a-helical secondary structure of Motif V (Fig. 2C). The other mostly conserved residues, Phe404, Val405, and Val406, interact primarily with other hydrophobic residues in the surrounding area. Generally, all of Motif V interacts with either a bound substrate or other highly conserved motifs that interact with bound substrate. Therefore, we investigated the importance of each residue in Motif V for viral replication.

We individually mutated every residue within NS3h Motif V in a luciferase-expressing WNV replicon system and determined replication efficiency for each mutant. Each Motif V mutation was designed to interrupt specific WT residue-residue interactions or residue-substrate interactions. As a negative control, Asp664 in the NS5 polymerase catalytic active site was mutated to valine (D664V) to disrupt viral genome replica-
tion (42–45). Additionally, the NS3 mutants A286L (Motif II) and R387M (Motif IVa) were used as controls for ablating ATPase activity and RNA-binding affinity, respectively (30, 46–50). The small hydrophobic side chain of Ala286 interacts with other hydrophobic residues stabilizing Motif II interactions with bound ATP. By introducing a bulky hydrophobic residue with A286L, the Motif II interactions with ATP will be perturbed, resulting in a disruption in ATP hydrolysis activity. The guanidium headgroup of Arg387 interacts with the phosphates of the RNA backbone. By removing the guanidium headgroup with R387M, important protein-RNA interactions will be interrupted. NS5 D664V and NS3 A286L and R387M ablated viral genome replication, as expected (Fig. 3A). All of the Motif V mutants, except for V405M, V406M, S411A, and A415G, ablated viral genome replication as well. V405M and S411A reduced replication to ~2.5%, whereas the I406M and A415G mutants reduce viral genome replication to less than 20%. T408S and S411A were 48% and 39% active, respectively, compared with WT NS3h. The results from the viral genome replication were projected onto the structure of Motif V (Fig. 3B). We noted that residues with ablated viral replication consisted mostly of the highly conserved residues, suggesting that these highly conserved residues that interact with either the RNA or the ATP play an important role in viral replication. Interestingly, Thr407 and Ala415, both highly conserved residues, still maintain a reduced level of viral replication, suggesting that there may not be a direct correlation between residue conservation and effect on replication.

As mentioned before, residues Thr407 and Ser411 interact with each other through a hydrogen bond (Fig. 2C). However, the T407A mutant nearly ablates genome replication, whereas the S411A mutation only reduces genome replication by ~60%. Both mutations should disrupt the hydrogen bond, but T407A and S411A have significantly different replication levels, suggesting that these two mutations have other interactions that potentially affect one or more of the NS3h functions during viral replication. These NS3h functions include the ability of the helicase to bind RNA, the ability to hydrolyze ATP, and the ability to unwind the dsRNA intermediate. Therefore, to determine whether or not these mutations have an effect on the individual helicase functions, we tested both T407A and S411A biochemically.

**NS3h residues Thr407 and Ser411 affect ATPase and helicase functions but not RNA-binding affinity**

We designed/adapted three biochemical assays to observe how the Thr407 and Ser411 mutations would affect the individual helicase functions. Recombinant WT, T407A, S411A, A286L (ATPase control), and R387M (RNA-binding control) NS3h were purified and tested in an RNA-binding affinity assay, an ATPase activity assay, and a helicase-unwinding activity assay. We used DENV4 NS3 proteins for this work due to established expression and purification protocols being in place (51) and the high degree of conservation between WNV and DENV in Motif V. To start, the RNA-binding affinity of WT, T407A, S411A, A286L, and R387M was measured in an RNA-binding affinity assay, an ATPase activity assay, and a helicase-unwinding activity assay. We used DEN45h NS5 proteins for this work due to established expression and purification protocols being in place (51) and the high degree of conservation between WNV and DENV in Motif V. To start, the RNA-binding affinity of WT, T407A, S411A, A286L, and R387M were tested via a fluorescence polarization assay (52). Fluorescence polarization for a single-stranded RNA oligonucleotide labeled with an Alexa 488 fluorophore (5 nM) was measured as the NS3 helicase increased in concentration (Fig. 4A). The binding affinity or $K_a$ for WT...
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Figure 3. Mutations in Motif V negatively affect viral genome replication. A, mutations to remove specific interactions within the central location of Motif V between ATPase active site and the RNA-binding cleft were tested in viral genome replication. B, results from the viral genome replication were projected onto the structure of Motif V with the hydrogen bond highlighted between residues Thr^{407} and Ser^{411}. The color bar ranges linearly from red (0% replication) to blue (100% replication). These data suggest that the hydrogen bond between Thr^{407} and Ser^{411} is important for viral genome replication. Error bars, S.E.

Figure 4. Experimental design of biochemical assays. A, RNA-binding affinity of NS3h was measured using fluorescence polarization of a single-stranded RNA (52). WT and NS3h variants are shown. B, ATPase activity was measured using BIOMOL Green absorbance at 650 nm over time. WT NS3h ATPase activity at five concentrations of ATP is shown here. C, RNA helicase-unwinding activity was measured by the increase of fluorescence over time. The WT NS3h data presented here are representative of the ATP-dependent unwinding activity obtained for each NS3h variant. D, WT, A286L, R387M, T407A, and S411A NS3h-unwinding activities were compared at 400 μM ATP. Error bars, S.E.

NS3h was 1.9 ± 0.8 μM, whereas the $K_d$ for R387M was 76.4 ± 31.0 μM (Table 1A). The R387M $K_d$ was significantly different compared with WT, suggesting that R387M does not bind the ssRNA as well as WT. Additionally, a similar trend was observed for WT and A286L. On the other hand, T407A (2.0 ± 1.1 μM) and S411A (1.4 ± 0.2 μM) were not significantly different as compared with WT. These data suggest that neither the T407A nor S411A mutation affected ssRNA binding to NS3.

We next investigated how these mutations affected NS3 ATPase activity. We utilized a colorimetric assay to observe ATP hydrolysis via Pi production and release for WT, T407A, S411A, A286L, and R387M (51). All reactions contained 50 nM ATP.
NS3, 5 nM dsRNA, and increasing concentrations of ATP. Progress curves were obtained for each NS3h at various ATP concentrations (Fig. 4B). We noticed at the high concentrations of ATP a substrate inhibition over time. Therefore, we used the substrate inhibition equation to calculate the enzyme kinetics for the ATPase activity assay. The catalytic efficiency ($k_{cat}/K_m$) for A286L (0.0033 ± 0.0022 × 10^-3 μM^-1 s^-1) was significantly reduced as compared with WT (0.0672 ± 0.019 × 10^-3 μM^-1 s^-1). The 20-fold decrease in catalytic efficiency was due to a significant increase in the $K_m$ for A286L (285.1 ± 150.5 μM) as compared with WT (17.3 ± 4.6 μM), whereas the turnover rate ($k_{cat}$) was consistent between the two systems (Table 1B). These data demonstrate that Ala^286 is critical for ATP hydrolysis. Additionally, the $k_{cat}/K_m$ for R387M (0.0314 ± 0.015 × 10^-3 μM^-1 s^-1) was reduced by a factor of 2 as compared with WT. Similar to A286L, the slight decrease in the catalytic efficiency of R387M was due to an increase in $K_m$ (49.5 ± 21.2 μM), whereas the $k_{cat}$ was not significantly different from WT. Interestingly, the increase in ATPase $K_m$ coupled with the decrease in RNA binding $K_d$ for R387M as compared with WT suggests that RNA binding promotes ATP binding, consistent with a previous report that ATP hydrolysis is stimulated by RNA (26). When WT NS3 helicase is compared with T407A and S411A, we observe similar turnover rates yet increased Michaelis–Menten constants. These results suggest that T407A and S411A hydrolyze ATP at a similar rate to WT, but more substrate is needed for the same catalytic efficiency. Overall, however, ATP hydrolysis was not increased or decreased by the T407A or S411A mutations.

Because both RNA binding and ATP hydrolysis were minimally affected by T407A and S411A, we then examined how these mutations affected overall helicase-unwinding activity. We utilized a molecular beacon–based helicase activity assay to identify the effects of translocation and unwinding of dsRNA for WT, A286L, R387M, T407A, and S411A (51, 53). Progress curves were obtained for each reaction containing NS3h variants (50 nM), dsRNA (5 nM), and increasing concentrations of ATP from the absence of ATP to 1 mM ATP (Fig. 4C). Each reaction was preincubated before the addition of ATP. For initial comparison, progress curves of WT, A286L, R387M, T407A, and S411A were normalized to the maximum relative fluorescence unit of WT, resulting in a relative unwinding activity compared with WT (Fig. 4D). The A286L (ATPase-defective) and R387M (RNA binding–defective) mutations ablated helicase-unwinding activity, as expected. The progress curves for T407A and S411A indicate that the mutations deviate from WT activity. To quantify the enzyme kinetics for each NS3h variant, we utilized a nonlinear regression fit on the progress curves that were not normalized. From the nonlinear regression, we determined an initial rate of reaction ($k_{obs}$) for each NS3h variant at all ATP concentrations. A plot consisting of $k_{obs}$ versus ATP concentration was fit to the Michaelis–Menten equation to determine the $k_{cat}$, $K_m$ and $k_{cat}/K_m$ (Table 1C). WT NS3 helicase unwound dsRNA with a $k_{cat}$ of 47.6 ± 4.4 × 10^-3 s^-1 and $K_m$ of 24.5 ± 11.3 μM (Table 1C). The dsRNA was unwound at a $k_{cat}$ of 80.8 ± 6.0 × 10^-3 s^-1 and a $K_m$ of 14.4 ± 6.2 μM for the T407A mutant, and at a $k_{cat}$ of 166.6 ± 15.5 × 10^-3 s^-1 and a $K_m$ of 58.0 ± 21.6 μM for S411A. These kinetics for T407A and S411A were significantly different compared with WT, which suggests that both mutations are catalytically more efficient than WT NS3h.

The interaction between Thr^407 and Ser^411 controls NS3 helicase function

T407A and S411A exhibit increased helicase activity, so we wanted to investigate if this was due to the absence or presence of a hydrogen bond between the two residues. Therefore, we constructed three additional mutations (T407C, S411C, and a...
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Figure 5. Examining the role of Thr<sup>407</sup> and Ser<sup>411</sup> interaction in NS3 helicase function. Additional mutations to force covalent bonds between the Thr<sup>407</sup> and Ser<sup>411</sup> residues were tested in viral genome replication. Error bars, S.E.

double mutant T407C/S411C) in both the replicon and recombinant protein systems to test viral genome replication, RNA-binding affinity, ATPase activity, and helicase-unwinding activity. Replacing residues Thr<sup>407</sup> and Ser<sup>411</sup> with cysteine residues would largely retain the hydrogen-bonding potential for each residue but would remove the methyl group from Thr<sup>407</sup>. T407C/S411C can be used to create a disulfide bond (∼2 Å) under oxidized conditions, locking 407 and 411 into a closer and more rigid interaction than a hydrogen bond (2.5–3 Å). First, we investigated the effects of T407C, S411C, and T407C/S411C on viral genome replication in our replicon system. The viral genome replication of T407C was significantly reduced (13.0 ± 8.0%), whereas the double mutant indicated a reduction to genome replication at 52.7 ± 30.0% as compared with WT NS3 helicase (Fig. 5). On the other hand, S411C exhibited genome replication activity similar to that of WT. These results suggest that the hydroxyl group of Thr<sup>407</sup> is responsible for part of the effect on genome replication because the cysteine mutation recovers replication by ∼2–3-fold compared with T407A. Additionally, these results indicate that the Thr<sup>407</sup> methyl group may be the important component of the threonine side chain due to the absence of viral replication in both T407A and T407C, which both lack the methyl group. The S411C mutant shows a complete recovery of replication efficiency, so the effect seen in S411A is due to the removal of the hydroxyl group. These data suggest overall that the hydrogen bond between Thr<sup>407</sup> and Ser<sup>411</sup> plays some role in replication, but the methyl in Thr<sup>407</sup> plays a different and potentially more important role in replication. Therefore, we investigated the individual NS3 helicase functions for these mutations in the recombinant protein system to further understand the role of the methyl and hydroxyl groups found in Thr<sup>407</sup> and Ser<sup>411</sup>. The T407C, S411C, and T407C/S411C mutants were expressed in BL21 DE3 pLysS competent cells to promote a reducing environment. WT and T407C/S411C mutant were also expressed in T7 Shuffle competent cells to promote an oxidizing environment for disulfide bond formation. All of the purified NS3 helicase variants were then tested in the three biochemical assays: RNA-binding affinity, ATPase activity, and helicase-unwinding activity. These cysteine mutations were tested under reducing conditions (TCEP) with the exception of the oxidized WT and T407C/S411C expressed in T7 Shuffle, which were tested in both reducing (TCEP) and nonreducing conditions. In all three assays, the BL21-expressed WT compared with T7 Shuffle—expressed WT have differing $K_p$, $k_{cat}$, and $K_m$ values, suggesting that the expression cell lines provide a different environment and therefore the T7 Shuffle—expressed NS3 helicases cannot be directly compared with the BL21-expressed helicases (Table 2). In the RNA-binding affinity assay, the BL21-expressed S411C and T407C/S411C were not significantly different in their binding affinities compared with WT, whereas T407C indicated a weaker RNA-binding affinity of 8.0 ± 2.5 μm, which was a 4.2-fold difference compared with WT (Table 2A). Additionally, the reduced T7-expressed T407C/S411C showed a 1.3-fold increase in $K_p$ as compared with the reduced T7 Shuffle—expressed WT. Similarly, the WT and T407C/S411C expressed in T7 Shuffle cells under nonreducing conditions were also not significantly different, with a 0.6-fold difference. All of these data suggest that T407C slightly decreases RNA-binding affinity potentially due to the lack of the methyl group, which may play a role in RNA binding to the helicase.

The NS3 helicase mutants expressed in BL21 and T7 Shuffle competent cells were tested for their ability to hydrolyze ATP. T7 Shuffle—expressed NS3 helicases were tested in both oxidizing and reducing conditions. T407C, S411C, and T407C/S411C expressed in BL21 cells all showed insignificant differences in both $k_{cat}$ and $K_m$ as compared with WT. The $k_{cat}$ for oxidized WT and T407C/S411C were not significantly different, with a $k_{cat}$ of 2.9 ± 0.8 × 10<sup>-3</sup> s<sup>-1</sup> and a $k_{cat}$ of 1.7 ± 0.2 × 10<sup>-3</sup> s<sup>-1</sup>, respectively, whereas the $K_m$ for WT (116.5 ± 49.7 μm) and T407C/S411C (49.9 ± 12.1 μm) under oxidized conditions were significantly different (Table 2B), suggesting that the disulfide bond present in the oxidized form of T407C/S411C increases ATP hydrolysis activity. On the other hand, WT and T407C/S411C expressed in T7 Shuffle cells under reduced conditions exhibit an insignificant difference for either $k_{cat}$ or $K_m$. Overall, these data suggest that hydrolysis of ATP is similar between WT and the cysteine mutations, meaning that the cysteine residues do not influence ATP hydrolysis directly.

Finally, the various cysteine NS3 helicase mutants as well as WT expressed in both BL21 and T7 Shuffle cells were tested in the helicase-unwinding activity assay. When comparing T407C, S411C, and T407C/S411C expressed in BL21 cells with WT, we observed a higher $k_{cat}$ with a lower $K_m$ for each mutant except for T407C, which exhibited a higher $K_m$ value (Table 2C). These data suggest that both S411C and T407C/S411C unwind dsRNA faster than WT although they do not bind ssRNA as well. T407C also unwinds dsRNA faster than WT, but the affinity for ssRNA is stronger, suggesting that T407C is a more catalytically efficient helicase overall compared with WT. The catalytic efficiency of WT and T407C/S411C expressed in T7 cells indicates no significant difference in the ability to unwind dsRNA substrates, suggesting that the double mutant exhibits similar overall unwinding activity compared with WT.
In silico mutations of Thr<sup>407</sup> and Ser<sup>411</sup> validate experimental assays

We further investigated how each mutant, alanine and cysteine, affected either the binding energy of the substrates or the overall secondary structure of NS3 helicase in the ssRNA + ATP substrate state using all-atom molecular dynamics simulations. Each mutant (A286L, R387M, T407A, T407C, S411A, S411C, and T407C/S411C) was simulated in triplicate for 1 μs using the ff14SB force field within the AMBER18 software package. Once the simulations were completed, the root mean square deviation was calculated in reference to the initial starting structure to determine the equilibration time that would be excluded for all other analyses. The three analyses performed on the simulations were a nonbonding interaction energy analysis, a probability of finding a lytic water in the ATP-binding pocket, and a projected covariance magnitude analysis. These analyses were specifically designed to compare results between simulation and the three biochemical assays.

First, we utilized the nonbonding interaction energy analysis to provide insight into how the mutations affect RNA-binding affinity from simulation. The nonbonding interaction energy was calculated between the bound ssRNA and the entire protein for each NS3 helicase variant. The more negative the linear interaction energy, the more strongly bound the ssRNA substrate is in the helicase RNA-binding cleft. All of the mutations exhibit strong ssRNA binding as compared with WT with the exception of A286L, R387M, and T407C mutants (Table 3A). The A286L and R387M mutations increase in their nonbonding interaction energies as compared with WT by 5.3 and 13.8%, respectively. These data suggest that A286L and R387M bind ssRNA weakly compared with WT. Additionally, T407C exhibits a nonbonding interaction energy of −7086.5 ± 36.7 kcal/mol. The more positive nonbonding interaction energy suggests that T407C binds ssRNA more weakly than WT. The lower nonbonding interaction energy correlates well with the reduced RNA binding affinity for T407C (Table 2A), validating our computational approach. Additionally, we observed an increase in side-chain mobility in the T407C mutant simulation compared with WT (Fig. S1), suggesting that the threonine at position 407 may control fluctuations within Motif V, allowing for the helicase to bind ssRNA strongly.

Next, we investigated how the mutations affect the ATP-binding pocket through determining the probability of finding a lytic water in the ATP hydrolysis active site. In order to do this, we utilized three collective variables to select for lytic waters: 1) the nucleophilic attack distance between the water oxygen atom and the ATP-terminal phosphoanhydride bond, 2) the nucleophilic attack angle between the water oxygen atom and the terminal phosphoanhydride bond of ATP, and 3) the angle between the water dipole moment and the terminal phosphoanhydride bond of ATP. Once the lytic waters were determined, the simulations were analyzed to determine the proba-
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**Table 3**

Thr<sup>407</sup> and Ser<sup>411</sup> mutations do not affect nonbonding interaction energies of bound ssRNA to NS3 helicase

| NS3<sub>h</sub> variant | Linear interaction energy (kcal/mol) | -Fold difference |
|------------------------|-------------------------------------|-----------------|
| WT                     | -747.4 ± 3.9                        |                 |
| A286L                  | -707.7 ± 0.0                        |                 |
| R387M                  | -644.5 ± 0.0                        |                 |
| T407A                  | -744.8 ± 23.0                       |                 |
| T407C                  | -708.6 ± 36.7                       |                 |
| S411A                  | -743.3 ± 11.9                       |                 |
| S411C                  | -736.7 ± 11.7                       |                 |
| T407C/S411C reduced    | -730.6 ± 27.2                       |                 |
| T407C/S411C oxidized   | -744.6 ± 18.4                       |                 |

A. RNA-binding energy

B. Projected covariance between Gly<sup>414</sup> and the helical gate

| NS3<sub>h</sub> variant | Response magnitude (Å²) | -Fold difference |
|------------------------|-------------------------|-----------------|
| WT                     | 0.032 ± 0.008           | 1.0             |
| A286L                  | 0.023 ± 0.0             | 0.7             |
| R387M                  | 0.060 ± 0.0             | 1.9             |
| T407A                  | 0.224 ± 0.088           | 7.0             |
| T407C                  | 0.045 ± 0.010           | 1.4             |
| S411A                  | 0.049 ± 0.007           | 1.5             |
| S411C                  | 0.057 ± 0.023           | 3.8             |
| T407C/S411C reduced    | 0.019 ± 0.001           | 0.6             |
| T407C/S411C oxidized   | 0.062 ± 0.021           | 1.9             |

The WT simulations indicated a projected covariance of 0.032 ± 0.008 Å², suggesting that there was a correlation between the fluctuations seen between the ATP-binding pocket and the helical gate. Both T407C (0.045 ± 0.010 Å²) and S411C (0.057 ± 0.023 Å²) were not significantly different compared with WT, although their -fold differences indicated an increase in projected covariance along the helical access. All of the rest of the mutations were significantly different compared with WT (Table 3B). Both A286L (0.023 ± 0.0 Å²) and reduced T407C/S411C (0.019 ± 0.001 Å²) showed a decrease in -fold difference (0.7 and 0.6, respectively) compared with WT, suggesting that these mutations negatively affect the dynamics of the helical gate (Table 3B). R387M, T407A, S411A, and oxidized T407C/S411C indicate an increase in -fold difference (1.9, 7.0, 1.5, and 1.9, respectively) compared with WT (Table 3B), suggesting that the fluctuations through Motif V positively affect the motion of the helical gate promoting translocation and unwinding of the dsRNA intermediate. Interestingly, when we compared the projected covariance magnitude -fold differences with the turnover rate -fold differences of the helicase-unwinding activity assay, all of the data supported each other except for A286L, R387M, and oxidized T407C/S411C. Both A286L and R387M were unable to unwind dsRNA in the helicase-unwinding assay, but in simulation, these mutants indicated a decrease and an increase in the projected covariance along the helical gate axis, respectively. The difference observed between simulation and experiment may be due to the pre-bound state of the simulation, whereas in experiment, the helicase still needs to bind the RNA. Additionally, the turnover rate of the oxidized double mutant indicated a decreased rate compared with WT, whereas the projected covariance magnitude indicated an increased fluctuation projected onto the helical axis compared with WT. This discrepancy might be due to the inability to control the state of the disulfide bond in experiment.

**Discussion**

In this study, a combination of computational, biochemical, and virological experiments were utilized to investigate the role of Motif V in NS3 helicase functions. WT simulations pointed to Motif V as a key player in the communication between the ATP-binding pocket and the RNA-binding cleft. To determine how Motif V may play a role in the communication, mutations
were introduced in both replicon and recombinant protein systems. The mutations were tested in a viral genome replication assay, and results suggested that the residues within Motif V affect how the virus replicates. From the viral replication assay, residues Thr\(^{407}\) and Ser\(^{411}\) sparked our interest due to the differing replication activity with prior knowledge of their interaction through a hydrogen bond that may stabilize Motif V structure. We tested both alanine and cysteine mutations in an RNA-binding affinity assay, an ATP hydrolysis activity assay, and a helicase-unwinding activity assay as well as simulated theses mutations in all-atom molecular dynamics. Results in this paper suggest that the methyl group in the side chain of Thr\(^{407}\) may play a role in decreasing helicase-unwinding activity through stabilizing interactions with residues Thr\(^{408}\), Lys\(^{366}\), and Arg\(^{387}\). These residues all interact with the bound RNA, allowing for viral replication. We can directly compare between the helicase-unwinding assay and viral replication assay because the helicase rates we observed biochemically appear to be in line with how fast dsRNA unwinding would likely need to occur in vivo during genome replication. Overall, our data suggest that Motif V is critical for the communication between the ATP-binding pocket and the RNA-binding cleft in NS3 helicase. Initial analyses of WT simulations supported findings reported by Davidson et al. (24) that Motif V may play a critical role in the communication between the ATP-binding pocket and the RNA-binding cleft. This led us to mutating all of the Motif V residues (residues 404–416) in the replicon system. Interestingly, only two Motif V mutations have been studied previously within the Flaviviridae virus family: T411A in HCV and G414A in dengue 2 virus (30, 54). Both of these mutations were tested biochemically but were not tested for their effect on viral genome replication. Therefore, our replicon data represent the first replication-based examination of the effect of Motif V mutations on viral genome replication. We observed that the majority of the mutations ablated viral replication activity with the exception of residues Val\(^{405}\), Val\(^{406}\), Thr\(^{407}\), Thr\(^{409}\), Ser\(^{411}\), and Ala\(^{415}\). Interestingly, most of the highly conserved residues ablated viral replication, but not all did, suggesting that the highly conserved residues can have some variability with the type of amino acid at those positions. Of the residues that did not ablate activity, Thr\(^{407}\) and Ser\(^{411}\) were of most interest due to the variable nature of these residues across all flaviviruses and the hydrogen bond that may stabilize the secondary structure of Motif V during replication. Therefore, we focus on these two positions for the remainder of the discussion.

Thr\(^{407}\) and Ser\(^{411}\) were mutated to alanine residues in the recombinant NS3 helicase and were expressed, purified, and tested in RNA-binding affinity, ATPase activity, and helicase-unwinding activity assays along with two controls, A286L and R387M. These three assays provide specific insight into how the helicase binds ssRNA and the rates of 1) ATP hydrolysis and 2) dsRNA unwinding. The rates determined from the latter two assays cannot be directly compared because the time scales are different because the ATPase assay is an end-point assay, whereas the helicase-unwinding activity assay is a continuous assay. The results from the RNA-binding affinity assay indicated that T407A and S411A bind ssRNA as strongly as WT while in the absence of ATP. We noted that ATP inhibited RNA binding in the fluorescence polarization assay, and thus ATP was left out of the reactions. The results from the ATP hydrolysis activity assay indicated that T407A and S411A hydrolyze ATP as well as WT. On the other hand, T407A and S411A seem to unwind dsRNA more quickly than WT in the molecular beacon helicase assay, suggesting that Thr\(^{407}\) and Ser\(^{411}\) are involved in the linkage between the ATP-binding pocket and the RNA-binding cleft. If the individual activities of RNA binding and ATP hydrolysis are unaffected by these mutations but the overall ability for the helicase to unwind dsRNA is affected, then these two mutations must play a role in the communication between the substrate-binding pockets. The observation that Thr\(^{407}\) and Ser\(^{411}\) increased the helicase turnover rate was surprising and suggested that one or both of these residues may naturally act to weaken the effect of ATP hydrolysis on helicase activity, giving rise to the intriguing possibility that the helicase has evolved to slow down its ATPase-dependent helicase activity. Position 407 is found predominately as a threonine, and 411 is found as serine (Fig. 2A and Table S1), but in a small group of
flaviviruses, position 407 can be alanine or serine and 411 can be alanine. This may indicate that the helicase function in other flaviviruses (yellow fever virus, Sepik virus, Entebbe bat virus, cell-fusing agent, Kamiti River virus, and Culex flavivirus) could be faster than most other flaviviruses.

We further investigated which aspects of the threonine and serine residues may play a critical role in the regulation of helicase-unwinding activity by introducing cysteines at positions 407 and 411 individually or as a double-mutant, T407C/S411C. T407C, S411C, and T407C/S411C were mutated in both the replicon and recombinant protein systems. We observed an increase in replication activity for the T407C mutation, a full recovery for the S411C mutation, and an increase in replication activity for the double mutant. T407C was the only mutation that weakened the ability of the helicase to bind ssRNA. The ATP hydrolysis activity is unchanged for all of the cysteine mutations, and the helicase-unwinding activity indicates that T407C and S411C increase unwinding activity, whereas the double mutant under oxidizing and reducing conditions remains as active as WT. The combination of these data suggests that the methyl group in the side chain of Thr^{407} may be important for slowing down helicase function. We observed a catalytically more efficient T407A in unwinding dsRNA, but a significant reduction in viral replication activity. When we mutate Thr^{407} to a cysteine (similar to a threonine but lacking in the methyl group), we recover some replication activity. When we mutated Ser^{411} to cysteine, we completely recover viral replication, suggesting that whereas the hydrogen bond between Thr^{407} and Ser^{411} is important, Thr^{407} has a separate function related to its methyl group. The methyl group in the Thr^{407} side chain normally interacts with the phenylalanine ring of Phe^{361} via hydrophobic interactions, potentially stabilizing the residues Thr^{408}, Lys^{366}, and Arg^{387}, which all interact with the RNA phosphate backbone (Fig. S1A). By removing the methyl from Thr^{407} with the T407C mutation, the side chain of position 407 is able to rotate and interact not only with Phe^{361} but also Leu^{385}, potentially causing RNA-interacting residues to become more variable with their interactions with bound ssRNA. As a result, these three residues that interact with the RNA may become more catalytically efficient in unwinding the dsRNA substrate. Results reported by Appleby et al. (37) examining residues equivalent to Arg^{387} and Thr^{408} (Arg^{393} and Thr^{411}) suggest that both residues play a direct role in the translocation along the ssRNA through conformational changes that are observed between nucleotide-bound and nucleotide-free states of HCV NS3. When comparing our results with HCV NS3, mutations at position 407 may disrupt the interactions that Arg^{387} and Thr^{408} have with the phosphodiester backbone of ssRNA such that Motif V does not undergo the same structural rearrangements to allow for Thr^{408} to shift to the next nucleotide in the bound ssRNA molecule as described previously (37). In the T407C mutation, the viral replication recovers some activity, suggesting that the hydroxyl group in Thr^{407} is also potentially important for proper viral replication due to the ability to form a hydrogen bond-like interaction with Ser^{411}. Overall, when the helicase becomes catalytically more efficient in the absence of the Thr^{407} methyl, viral genome replication decreases. It is unclear why this is the case, but one hypothesis is that reduced replication may be due to a rate mismatch between T407A, T407C, and S411A NS3 and the RNA-dependent RNA polymerase of NS5. If this is the case, it may suggest that the NS3 helicase and NS5 polymerase have co-evolved to match their catalytic rates so that RNA is fed at an optimal rate between the two enzymes.

All of the NS3 helicase mutations were simulated using the ff14SB force field within the AMBER18 software package to better define how the mutations affected NS3 helicase function. Three analyses were performed on the simulations to determine how the structure of the helicase has changed due to the presence of the mutations. The first analysis we utilized was the nonbonding interaction energy analysis. The interaction energies determined from this analysis were compared with the RNA-binding affinity ($K_d$) obtained from experiment. We observed that in both computation and experiment, the mutations (T407A, T407C, S411A, S411C, and T407C/S411C) did not significantly change the ability of the helicase to bind ssRNA. The only mutation that weakened the binding of ssRNA in the helicase was R387M, which was expected because this mutation was a control for RNA binding. As we moved forward with our analysis of the simulations, we investigated how the mutations affected the probability of finding a lytic water in the ATPase active site (Fig. 6). When we compared the probability of finding a lytic water in the ATP-binding pocket with the turnover rate determined from the ATPase assay, we observed that none of the mutations significantly changed the probability of finding a lytic water. When comparing the probability of a lytic water with the turnover rate of the helicase activity, we observed that S411A and S411C were set apart from the rest of the mutations, suggesting that these two mutations play a role in controlling the rate of unwinding activity through the hydrogen bond with Thr^{407}. Interestingly, when Thr^{407} is mutated to a cysteine, we observe an increase in mobility at position 407 (Fig. S1). The cysteine residue lacks the methyl group and contains a thiol group instead of a hydroxyl group found in the threonine side chain. The lack of the methyl group seems to allow for the thiol group to fluctuate frequently throughout the simulation (Fig. S1, B and C), suggesting a potential hydrophobic interaction between Thr^{407} and nearby hydrophobic residues (such as Leu^{385}) that may influence the position of Thr^{407} within Motif V. This supports our previous results for Thr^{407} in that the orientation of the methyl groups is critical for helicase function. Additionally, the projected covariance magnitude analysis indicated that T407A increased by 7.0 compared with WT, suggesting that the methyl and hydroxyl group of threonine are important for stabilizing the interactions of position 407 with its surrounding residues. Without either of those functional groups at position 407, Motif V may allow more structural changes to occur that positively affect the helical gate. This is evident when we take into account the projected covariance of T407C. T407C indicated a slight increase compared with WT, which is similar to what is observed in the $k_{cat}$ of T407C from the helicase-unwinding assay. The thiol group of the cysteine potentially interacts with Ser^{411}, stabilizing Motif V enough to control the structural changes within the helicase to allow for optimal translocation and unwinding of dsRNA.
This work provides detailed computational, biochemical, and virological insight into how the helicase utilizes the energy produced from ATP hydrolysis to power translocation and unwinding of the viral dsRNA intermediates. We have learned that the methyl group in Thr\textsuperscript{407} is potentially critical for regulating the helicase to maintain an optimal rate of translocation and unwinding during replication. If the Thr\textsuperscript{407} is mutated to an alanine or a cysteine, the viruses we examined are not able to replicate the viral genome efficiently \textit{in vitro}. This Thr\textsuperscript{407} residue within Motif V interacts with Ser\textsuperscript{411}, which seems to properly orient Thr\textsuperscript{407}, potentially stabilizing the structure of Motif V. When we remove that interaction between Thr\textsuperscript{407} and Ser\textsuperscript{411} with a mutation to Ser\textsuperscript{411}, the helicase is able to unwind dsRNA intermediates with more catalytic efficiency, suggesting that Thr\textsuperscript{407} is responsible for regulating translocation and unwinding dsRNA intermediates. As previously mentioned, the role of Motif V in SF2 helicases is not well-understood. This work has provided computationally driven experimental insight into how Motif V plays a role in the communication between the ATP-binding pocket and the RNA-binding cleft in the viral DEAH subfamily of SF2 helicases \cite{6}. Additional work will be required to determine whether the results we observe here for flaviviruses hold in other subfamilies of SF2 helicases. Ultimately, the data from this paper suggest that flaviviruses may utilize suboptimal NS3 helicase activity for optimal genome replication and that Motif V may play an inhibitory role for helicase activity during viral infection \textit{in vitro} and \textit{in vivo}. The role of Motif V in controlling helicase function will need to be further defined, which may eventually lead to the development of novel vaccine candidates or antiviral drug targets.

**Experimental procedures**

**Plasmids and mutagenesis**

The DNA expression plasmid used in the protein purification protocol for dengue 4 NS3 (accession number AQV12689) was adapted from Luo \textit{et al.} \cite{6}. The QuikChange mutagenesis (Agilent Technologies) method was used to mutate specific residues (A286L, R387M, T407A, T407C, S411A, S411C, and T407C/S411C) within the WT dengue NS3h plasmid optimized for expression in \textit{Escherichia coli}. Plasmids for each mutation were submitted to GENEWIZ for sequencing to verify each sequence.

**Protein expression and purification**

WT dengue NS3h gene with an N-terminal thioredoxin, His\textsubscript{6} tag, and thrombin cleavage site was synthesized and cloned into a T7 expression plasmid. The WT NS3h construct was transformed into BL21 DE3 pLysS or T7 Shuffle \textit{E. coli} competent cells. 5 ml of Luria broth cultures containing ampicillin (50 \textmu g/ml) and chloramphenicol (34 \textmu g/ml) were grown overnight at 37 °C. The next day, the entire 5-ml culture was added to a 750-ml Luria broth culture containing CaCl\textsubscript{2} and stored at 4 °C. A second nickel column was subsequently used to remove thioredoxin from NS3h, which was collected in the flow-through fraction. The flow-through fraction was concentrated using a Vivaspin Turbo 15 ultrafiltration spin column \cite{5} and then applied to a HiLoad 16/600 Superdex 200 pg (GE Healthcare) column on an AKTA Pure FPLC system \cite{6}. The protein complex was eluted off the nickel column with a linear gradient of high-imidazole buffer (HIB) (50 mM Tris-base, pH 8.00, 400 mM NaCl, 10 mM imidazole, and 5% glycerol). Fractions containing the NS3h complex were collected and dialyzed against 350 ml of LIB containing CaCl\textsubscript{2} for 2 h using a Slide-A-Lyzer dialysis cassette (3–12 ml (Life Technologies, Inc.)). 75 \textmu l of soluble thrombin (GE Life Sciences) was added to the dialyzed protein for an overnight incubation at 4 °C. A second nickel column was subsequently used to remove thioredoxin from NS3h, which was collected in the flow-through fraction. For the protein purification, LIB (20 ml) was added to the frozen bacterial pellets. Once thawed, bacterial cells were disrupted three times on a 110S Microfluidizer (Microfluidics), the lysate was clarified by centrifugation using a JA-25.50 rotor for 20 min at 17,000 rpm at 4 °C, and supernatant was filtered using a 0.45-\textmu m syringe filter. Protein purification was achieved using a multistep approach. The thioredoxin-dengue NS3h complex was first purified from the filtered lysate with nickel affinity chromatography using a HisTrap HP (GE Healthcare) column on an AKTA Pure FPLC system \cite{6}. The protein complex was eluted off the nickel column with a linear gradient of high-imidazole buffer (HIB) (50 mM Tris-base, pH 8.00, 400 mM NaCl, and 5% glycerol).

**RNA helicase–unwinding activity assay**

For the ATP-dependent helicase assay \cite{51, 53}, all reactions were performed in 96-well black microplates (Thermo Scientific) in a final volume of 120 \textmu l, and they contained 50 nM NS3h, 0.05 mM TCEP, 0.01% Tween 20, 5 \mu g/\mu l BSA, 1.25 mM MgCl\textsubscript{2}, 25 mM MOPS (pH 6.5), 5 mM RNA complex, and varying concentrations of ATP (Jena Bioscience) ranging from 1 \mu M to 1 mM. The RNA complex consists of an RNA-tagged Alexa 488 oligonucleotide (5’-AGUGCGCU-GUAUGCUGAAGG-3’-AlexF488N) and a DNA-tagged Iowa Black fluorescent quencher (5’-IABkFQ-CCTACGCACACAGCTCGTAGG-3’). The RNA template (5’-GGAGCUGUUGGCUGAAAGCAGACAGCUUAAAGG-3’). The RNA complex was annealed through a slow cooling process by boiling H\textsubscript{2}O to 95 °C and allowing it to cool to room temperature over 1 h. The control with no MgCl\textsubscript{2} contained 20 mM EDTA. Reactions were scanned individually at 37 °C using a Victor X5 multilabel plate reader (PerkinElmer Life Sciences) instrument for 300 s. Rates were determined from where the progress curves were exponential and linear with time. Data were then fit to the Michaelis–Menten equation (Equation 1) using Python2.7.
**NS3 Motif V acts as a regulator of energy transduction**

\[ v = \frac{V_{\text{max}}[S]}{(K_m + [S])} \]  

(Eq. 1)

where \( V_{\text{max}} = k_{\text{cat}}[E] \), \( v \) is the velocity, \( k_{\text{cat}} \) is the apparent first-order rate constant in \( s^{-1} \), \([E]\) is the concentration of the enzyme, \([S]\) is the concentration of the ATP substrate, and \( K_m \) is the concentration of the substrate at one-half \( k_{\text{cat}} \). The \( K_m \), turnover rate \( (k_{\text{cat}}/K_m) \), and specificity constant \( (k_{\text{cat}}/K_m) \) were determined from fitting to Equation 1.

**RNA ATPase activity assay**

A colorimetric ATPase activity assay was derived from the RNA helicase assay (51). The contents of the reactions remained the same. Using a 96-well clear plate, reactions were injected with BIOMOL Green (Enzo) and incubated at 37 °C for 15 s before measuring the absorbance at 650 nm on a Victor X5 multilabel plate reader. To obtain an entire curve for an individual ATP concentration, reactions were measured every 90 s for 12.5 min. Rates were determined from progress curves. Data were then fit to the substrate inhibition equation (Equation 2) using the R software environment (55),

\[ v = \frac{V_{\text{max}}[S]}{(K_m + [S])\left(1 + \frac{[S]}{K_i}\right)} \]  

(Eq. 2)

where \( K_i \) is the inhibition constant. From the fit, the \( k_{\text{cat}} \), \( K_m \), \( k_{\text{cat}}/K_m \), and \( K_i \) were determined for the ATPase assay. \( K_i \) data are reported in Table S2.

**RNA-binding affinity assay**

A fluorescence polarization assay was also adapted from the RNA helicase assay to determine RNA-binding \( K_d \) as described previously (52). The fluorescence polarization assay reactions contained increasing concentrations of NS3h, 0.05 mM TCEP, 0.01% Tween 20, 5 μg/μl BSA, 1.25 mM MgCl₂, 25 mM MOPS (pH 6.5), and 5 mM ppAGUAA-tagged Alexa 488 RNA oligonucleotide. All reactions were incubated at 37 °C for 10 min in a 384-well black plate within the plate reader before measuring the fluorescence polarization of the Alexa 488 fluorophore. Data were analyzed using nonlinear regression analysis in Prism8 using a maximum value of 400.0 based on protein binding reported previously (52).

**Cell culture and virus replicon plasmids**

Baby hamster kidney (BHK) cells were maintained in HyClone Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum, 50 μM HEPES (pH 7.5), 0.5% penicillin/streptomycin, and 5% l-glutamine. Cells were grown in humidified incubators at 37 °C with 5% CO₂. West Nile virus replicon utilized in the viral genome replication assay was designed as a plasmid-launched virus replicon expressing the firefly luciferase as described previously (56, 57). Additional site-directed mutagenesis was performed on the WT WNV replicon using the QuikChange mutagenesis (Agilent Technologies) method to produce the following mutations: NS5 D664V and NS3h A286L, R387M, F404K, V405M, V406M, T407A, T407C, T408S, D409A, I410G, S411A, S411C, E412D, M413A, G414A, A415G, N416D, and T407C/S411C double mutant. The presence of the mutations was confirmed through GENEWIZ sequencing.

**Viral genome replication assay**

BHK cells were plated into 24-well plates at 50,000 cells/well and allowed to adhere to the plates overnight. The next day, WT and mutant WNV replicon plasmids were transfected into the BHK cells at 250 ng/well using Lipofectamine2000 (Life Technologies). 48 h post-transfection, the cells were lysed with 1× lysis buffer from the Luciferase Assay Kit (Promega), and 20 μl of the lysates were transferred into a white 96-well plate. Using the plate reader, 100 μl of the luciferase assay reagent from the Luciferase Assay Kit was injected into each sample before measuring the luminescence of the firefly luciferase signal. Data were obtained with an n ≥ 3. The mean and S.E. were plotted using Matplotlib (58).

**All-atom molecular dynamics simulations**

An exemplar structure of the simulated DENV4 NS3h ssRNA + ATP substrate state (Protein Data Bank code 2JLV) was used for all simulations (24, 33). The ssRNA + ATP state was run for nine systems of NS3h (WT, A286L, R387M, T407A, T407C, S411A, S411C, and the double mutant T407C/S411C in oxidized and reduced conditions) in triplicate for 1 μs each. All of the mutations were generated from the WT NS3 helicase. All of the NS3h systems were simulated in explicit solvent MD using the AMBER18 software package: ff14SB (protein) and RNA.O1L3 (RNA) (59). Parameterization files for the ATP molecule (60) were obtained from the AMBER parameter database. Each system was solvated with a cubic box of TIP3P water with an average dimension of 85 Å. Sodium and chloride ions were added at a concentration of 100 mM to neutralize the system. The simulations were performed with periodic boundary conditions in an isothermal-isobaric (NPT) ensemble with a stochastic barostat of 1 bar and a Langevin thermostat of 310.0 K. The nonbonding interactions cutoff was 12 Å; particle mesh Ewald was used for treating long-range electrostatics; and hydrogens were constrained with the SHAKE algorithm. An integration time step of 2 fs was used. Positions and energies were written every 2 ps.

All systems were initially minimized with a 10,000-step minimization with harmonic restraints (force constant of 75 kcal mol⁻¹ Å⁻²) placed on all solute atoms and a 10,000-step minimization with no restraints. Following the minimization, each system was heated from 50.0 to 310.0 K in 10-K increments with a harmonic restraint of 75 kcal mol⁻¹ Å⁻² on all solute atoms. Last, a series of five equilibration steps were implemented to slowly remove the harmonic restraint.

**Data analysis of MD simulations**

All analyses of MD simulations were performed using Python 2.7 and the MDAnalysis package (61, 62). These analyses include the linear interaction energy analysis, the lytic water analysis, and the projected covariance magnitude analysis. Further details of each analysis, except for the linear interaction energy, are described below. Plots were created using Matplotlib (58), and VMD (9) was utilized to visualize simulations and MDKQ3F0Q5.
generate structural images. All scripts for the analyses are available on GitHub (https://github.com/mccullaghlab/T407_S411_Mutants_of_NS3h).

Waters within the NTPase active site were defined as “lytic” using three collective variables: 1) the nucleophilic attack distance between the water oxygen atom and the ATP γ-phosphorus atom, 2) the nucleophilic attack angle between the water oxygen atom and the terminal phosphoanhydride bond of ATP, and 3) the dipole moment angle between the water molecule’s dipole moment vector and the terminal phosphoanhydride bond of ATP. The first two metrics describe the geometric positioning of waters within the hydrolysis active site. Waters that have a nucleophilic attack distance less than 5 Å and a nucleophilic attack angle greater than 160° have the potential to be the lytic water due to their positioning relative to the terminal phosphate group. The third metric describes the chemically relevant orientation of a water in regard to its nucleophilic attack on this phosphate group. An ideally positioned water with a dipole moment angle of greater than 90° is defined as “lytic.” A water that meets all three of these collective variable conditions is positioned in a small volume of the NTPase active site, is hydrogen-bonding with the active site’s proton acceptor (Glu285) (data not shown), and the water’s dipole moment vector is approximately facing the terminal phosphoanhydride bond vector, as proposed in the SN2 mechanism for the hydrolysis reaction.

The projected covariance magnitude represents the fluctuation between Gly414 (source residue) and Ser364 and Lys366 (sink residues) projected onto the vector spanning the helical gate and large magnitude covariance with Gly414. The helical gate and large magnitude covariance with Gly414.

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