The C-terminal 50 Amino Acid Residues of Dengue NS3 Protein Are Important for NS3-NS5 Interaction and Viral Replication*

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Dengue virus multifunctional proteins NS3 protease/helicase and NS5 methyltransferase/RNA-dependent RNA polymerase form part of the viral replication complex and are involved in viral RNA genome synthesis, methylation of the 5' cap of viral genome, and polyprotein processing among other activities. Previous studies have shown that NS5 residue Lys-330 is required for interaction between NS3 and NS5. Here, we show by competitive NS3-NS5 interaction ELISA that the NS3 peptide spanning residues 566–585 disrupts NS3-NS5 interaction but not the null-peptide bearing the NS70A mutation. Small angle x-ray scattering study on NS3(172–618) helicase and covalently linked NS3(172–618):NS5(320–341) reveals a rigid and compact formation of the latter, indicating that peptide NS5(320–341) engages in specific and discrete interaction with NS3. Significantly, NS3:Asn-570 to alanine mutation introduced into an infectious DENV2 cDNA clone did not yield detectable virus by plaque assay even though intracellular double-stranded RNA was detected by immunofluorescence. Detection of increased negative-strand RNA synthesis by real time RT-PCR for the NS3:NS70A mutant suggests that NS3-NS5 interaction plays an important role in the balanced synthesis of positive- and negative-strand RNA for robust viral replication. Dengue virus infection has become a global concern, and the lack of safe vaccines or antiviral treatments urgently needs to be addressed. NS3 and NS5 are highly conserved among the four serotypes, and the protein sequence around the pinpointed amino acids from the NS3 and NS5 regions are also conserved.

Dengue virus (DENV),2 of which there are four distinct serotypes (DENV1–4), is an important re-emerging mosquito-borne flavivirus that is endemic in more than 100 countries, causing >390 million human infections that result in ~100 million dengue fever (DF) cases (1). Infections with DENV are either asymptomatic or can result in self-limiting febrile illness (DF) that leads to a broad spectrum of pathologies, including severe DF without hemorrhagic symptoms, dengue hemorrhagic fever, or dengue shock syndrome. The current treatment for dengue infection is mainly supportive, and there is no preventative vaccine or effective antiviral agents to treat DF or the more severe disease manifestations.

The ~11-kb positive-sense single-stranded RNA genome of DENV serotypes share around 70% sequence identity and contain a 5'-type-1 cap as well as 5'- and 3'-untranslated regions (UTR) that flank a single open reading frame (ORF) (2). An ~3300-amino acid polyprotein precursor is translated from the ORF and is processed by both host and viral proteases to yield three structural proteins (capsid, premembrane protein, and envelope protein (E)) and seven nonstructural proteins (NS) (NS1, NS2A, NS2B, NS3, NS4A, NS4B, and NS5) that are

1 The abbreviations used are: DENV, Dengue virus; SAKS, small-angle x-ray scattering; RuRP, RNA-dependent RNA polymerase; P/S, penicillin/streptomycin; bNLS, importin b-mediated nuclear localization sequence; NSD, normalized spatial discrepancy; IFA, immunofluorescence assay; EOM, ensemble optimization method; E, envelope protein; DF, dengue fever; RC, replication complex; Y2H, yeast two-hybrid.

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Background: NS3-NS5 interaction is important for the dengue virus life cycle.

Results: NS3 residue Asn-570 is essential for its interaction with NS5; mutation in an infectious cDNA abolished virus production and reduced positive-strand RNA synthesis.

Conclusion: NS3-NS5 interaction may be required for coordinated positive- and negative-strand RNA synthesis.

Significance: NS3-NS5 interaction may be a target for rational design of antiviral drugs.
involved in formation of mature virion and viral RNA replication, respectively. Among the NS proteins, NS3 and NS5 contain the enzymatic activities that are essential for DENV replication (3).

Both NS3 (residues 1–618) and NS5 (residues 1–900) are large multifunctional proteins with sequence identity of around 60 and 70% among the four serotypes, respectively (3). NS3 contains an N-terminal serine protease domain (residues 1–170) that requires NS2B to be an active protease (4–8). Its C-terminal domain contains ATPase/helicase activity for unwinding of the double-stranded RNA (dsRNA) intermediate (9–14) and RNA 5′-triphosphatase activity for viral RNA 5′-capping reaction that is carried out together with the N-terminal domain of NS5, which has methyltransferase activity (3, 9, 13, 15–22). The C-terminal domain of NS5 has RNA-dependent RNA polymerase (RdRP) activity, which is crucial for RNA replication (19, 23–26). NS3 and NS5 have been shown to interact and colocalize in infected cells, and NS3 RNA 5′-triphosphatase activity has been reported to be stimulated by NS5 in vitro (9, 27–29). These observations are consistent with the functional roles of both NS3 and NS5 in the replication complex (RC) (30, 31).

During viral RNA replication within the RC, many critical RNA-RNA, RNA-protein, and protein-protein interactions occur to synthesize both positive- and negative-strand viral RNA (31, 32). There have been several reports of NS3-NS5 interactions that include biochemical pulldown assays from infected cell extracts (28, 32–34) and two-hybrid (Y2H) studies that mapped the interaction to the C-terminal region of NS3 helicase (residues 303–618) and the N-terminal region of NS5 RdRP (residues 320–368; known as bNLS (nuclear localization sequence)) (35, 36). The NS5-binding site appears to be centered at residue Lys-330 because the mutation to alanine disrupted its interaction with NS3 and abolished RNA replication, although the in vitro RdRP activity was unaffected (37). Based on available crystal structures of the RdRP domain of NS5, it has been proposed that the cavity occupied by Lys-330 may be a potential target for antiviral drug design by blocking NS3-NS5 interaction (37). However, the details of the interaction from the NS3 perspective is missing to fully exploit structure-guided drug design.

In this study, through the use of both NS3 WT and mutant peptides in competitive NS3-NS5 interaction ELISA, we identified a conserved amino acid in subdomain III of DENV NS3 protein, Asn-570, as being critical for its interaction with NS5. Mutation of NS3:Asn-570 to alanine in the DENV2 cDNA clone abolished infectious virus production and reduced viral protein production and RNA replication. This mutation also suggests that the NS3-NS5 interaction is essential for viral RNA replication by possibly coordinating positive- and negative-strand synthesis. Small angle x-ray scattering (SAXS) data of NS3 helicase (residues 172–618) covalently linked to NS5(320–341) supports the observation that physical interaction occurs in the region of interaction between NS3 and NS5.

### EXPERIMENTAL PROCEDURES

**Cell Lines and Viruses**—Baby hamster kidney cells (BHK-21) were cultured in RPMI 1640 medium (Invitrogen) supplemented with 10% fetal bovine serum (FBS), 100 units/ml penicillin, and 100 μg/ml streptomycin (1% P/S) in 5% CO₂ at 37 °C.

Human hepatoma (Huh-7) cells were cultured in DMEM (Invitrogen) supplemented with 10% FBS and 1% P/S at 37 °C, with 5% CO₂. *Aedes albopictus* mosquito (C3/36) cells were cultured in RPMI 1640 medium supplemented with 25 mM HEPES, 10% FBS, and 1% P/S at 28 °C, in the absence of CO₂.

DENV2 of cosmopolitan genotype (GenBank™ number EU081177.1) that was used in this study was grown in C3/36 cells and titered in BHK-21 cells before storage at ~80 °C. This virus was isolated during a local dengue outbreak that occurred in 2005 as part of Early Dengue Infection and Outcome (EDEN) Study in Singapore (38).

**Plasmid Construction**—Plasmids for the expression of DENV3 NS5 RdRP (residues 273–900), NS3 full-length protein (NS2B18NS3; residues 49–66 of NS2B and 1–618 of NS3 connected by a flexible linker G₄S₄G₄), and DENV4 NS3 helicase domain (NS3(172–618)) have been described previously (Fig. 1A and Table 1) (11, 25, 39).

### TABLE 1

List of proteins/peptides used in ELISA, SAXS and viral inhibition assay

| Protein/peptide name | Vector | Tag | Residue number | Ref. |
|----------------------|--------|-----|----------------|-----|
| His-NS5 RdRP         | pET15b (Novagen) | His₆ (not removed) | 273–900 of DENV3 NS5 | Yap et al. (25) |
| NS2B₁₈NS3            | pET32b (Novagen) | Glutathione S-transferase (GST; not removed) | 49–66 of DENV3 NS2B and 1–618 of DENV3 NS3 connected by G₄S₄G₄ linker | Luo et al. (11) |
| His-NS3(172–618)     | pET14b (Novagen) | His₆ (not removed) | 172–618 of DENV3 NS3 | In this study |
| GST-NS3(482–618)     | pGEX-4T-1 (GE Healthcare) | GST (not removed) | 482–618 of DENV3 NS3 | In this study |
| His-NS3(566–585)     | pET32b (Novagen) | His₆ (removed) | 566–618 of DENV3 NS3 | Luo et al. (11) |
| His-NS3(566–585)     | pET32b (Novagen) | His₆ (not removed) | 566–618 of DENV3 NS3 | In this study |
| NS3(570–585)         | Synthesized by NTU peptide synthesis core facility (Singapore) | | 570–585 of DENV3 NS3 | In this study |
| NS3(86–100)          | Biotin | Third α-helix of the Antennapedia homeodomain | 86–100 of DENV3 NS3 | (56) |
| Penetratin (p)       | pNS5(320–341) | | | 320–341 of DENV3 NS5 connected to penetratin sequence |
| Penetratin (p)       | pNS5(320–341) | | | Scrambled sequence of 320–341 of DENV3 NS5 connected to penetratin sequence |
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DENV3 NS3(172–618), NS3(482–618), and NS3(566–618) fragments were amplified from DENV3 NS2B18NS3 construct (as mentioned above). NS3(172–618) fragment was amplified with forward primer 5′-CACAAGATACGTTGACCTTCTGGTGGACCAACC-3′ and the reverse primer 5′-GTGCTCGAGTGGAGATCCAGCTTCTTTTGC-3′. The underlined sequence corresponds to NdeI and BamHI sites, respectively. The PCR fragment was digested with NdeI and BamHI and cloned into pET14b (Novagen, Germany). The NS3(482–618) fragment was amplified with the forward primer 5′-CCAGCTCTTCTGCGAGTAACTCC-3′ and the reverse primer 5′-CATGACCAGTGAAGACCATG-3′. Product was digested with NdeI and BamHI and cloned into pET14b (Novagen, Germany). The NS3(566–618) fragment was amplified with the forward primer 5′-GAAATGGTGCGGACAGTCAGG-3′ and the same reverse primer as above. The underlined sequence corresponds to BamHI and XhoI sites, respectively. Both PCR products were digested with BamHI and XhoI and cloned into pGEX-4T-1 (GE Healthcare) (Fig. 1A and Table 1).

DENV4 NS3(172–618)-NS5(320–341) fragment (connected by a flexible linker G4SG4, Fig. 3A and Table 1) was amplified from DENV4 NS2B18NS3 construct (as mentioned above). The full-length PCR product was generated by first amplifying the helicase region with the forward primer 5′-GACAACCAGCTTCTTCTGAGACCTCC-3′, followed by two separate PCRs that added the flexible linker sequence and NS5(320–341) sequence to the helicase via two reverse primers, 5′-GAAATGGTGCGGACAGTCAGG-3′ and the reverse primer 5′-CTCTTCTCCTCTGAGACCTCC-3′, and 5′-GGCGGCGCGACTGAGCTC-3′ and 5′-GGCGGCGCGACTGAGCTC-3′. The underlined sequence corresponds to the G4SG4 linker and NS5(320–341) sequence, respectively. The PCR product was digested with NdeI and SalI and cloned into pProEx HTb vector.

Mutation of NS3 Asn-570 to alanine in DENV3 NS2B18NS3 was done using QuikChange II XL site-directed mutagenesis kit (Stratagene), according to the manufacturer’s protocol. The following primers were used: NS3 N570A forward (5′-GATGGGC-3′) and reverse primer 5′-CACGAAGATACGTTGACCTTCTGGTGGACCAACC-3′ and the reverse primer 5′-CATGACCAGTGAAGACCATG-3′. The underlined sequence corresponds to the NdeI and BamHI sites, respectively. The PCR product was digested with NdeI and BamHI and cloned into pET14b (Novagen, Germany). The NS3(482–618) fragment was amplified with the forward primer 5′-CCAGCTCTTCTGCGAGTAACTCC-3′ and the reverse primer 5′-CATGACCAGTGAAGACCATG-3′. The underlined sequence corresponds to the NdeI and BamHI sites, respectively. The PCR product was digested with NdeI and BamHI and cloned into pET14b (Novagen, Germany). The NS3(566–618) fragment was amplified with the forward primer 5′-GAAATGGTGCGGACAGTCAGG-3′ and the reverse primer 5′-CTCTTCTCCTCTGAGACCTCC-3′ and the reverse primer 5′-CATGACCAGTGAAGACCATG-3′. The underlined sequence corresponds to the NdeI and BamHI sites, respectively. The PCR product was digested with NdeI and BamHI and cloned into pET14b (Novagen, Germany).

**Protein Expression and Purification**—For ELISA and ATPase activity assay, NS3 and NS5 constructs (Table 1) were transformed into Escherichia coli BL21 CodonPlus (DE3)-RIL cells (Stratagene) for protein expression and purified as previously published (11, 25, 39).

For SARS, NS3(172–618) and NS3(172–618)-NS5(320–341) constructs were transformed into E. coli BL21 CodonPlus (DE3)-RIL cells (Stratagene) for protein expression and purified as described below. The cells were disrupted on ice by sonication three times for 1 min in buffer A (20 mM sodium phosphate, pH 7.4, 500 mM NaCl, 0.8 mM DTT, and 2 mM PefablocSC (Biomol)). Precipitated material was separated by centrifugation at 12,500 × g for 25 min, and the supernatant was filtered (0.45 μm; Millipore). The filtered supernatant was incubated with 2 ml of nickel-nitrilotriacetic acid-agarose (Qiagen) for 1.5 h at 4 °C, and the His6-tagged protein was eluted with an imidazole gradient from 20 to 200 mM in buffer A. Fractions containing the protein of interest were pooled and subjected to overnight cleavage with thrombin for NS3(172–618) or tobacco etch virus protease for NS3(172–618)-NS5(320–341) and dialyzed overnight in buffer B (20 mM sodium phosphate, pH 7.4, 200 mM NaCl) at 4 °C. Following incubation, the dialyzed sample was incubated with 1 ml of nickel-nitrilotriacetic acid-agarose for 1 h at 4 °C, and the flow-through containing cleaved NS3(172–618) or NS3(172–618)-NS5(320–341) was collected and applied to a gel filtration column (Superdex™ 200 HR 10/300 column, GE Healthcare) in buffer C (20 mM Tris/HCl, pH 7.4, 200 mM NaCl, 5% glycerol, 1 mM DTT). Fractions containing NS3(172–618) or NS3(172–618)-NS5(320–341) were pooled and concentrated using Amicon Ultra-4 centrifugal unit (10-kDa molecular mass cutoff; Millipore).

**Peptide Synthesis**—Peptides were synthesized at the Nanyang Technological University peptide synthesis core facility (Singapore).

**Competitive NS3-NS5 Interaction ELISA**—The ELISA (Fig. 1B) was performed as published previously (33). The concentration of NS2B18NS3 was fixed at 60 or 80 nM and mixed with increasing concentrations of truncated NS3 proteins (DENV3 NS3(172–618), GST-NS3(482–618), GST-NS3(566–618), or GST (negative control)) or NS3 peptides (NS3(566–585) and NS3(576–585)) (NS50A), NS5(571–585), or NS86 (100 ng/mL) (negative control); listed in Table 1). The concentration of NS3 protein at which 50% inhibition of NS2B18NS3 protein binding occurs represents the apparent KD value for the DENV3 NS3 RdRP-NS3 protein interaction. Data were fitted to the sigmoidal dose-response equation (variable slope) by nonlinear regression using GraphPad Prism 5 from triplicate measurements.

**Viral Inhibition Assay**—2 × 10⁵ Huh-7 cells were seeded into a 12-well plate and incubated overnight at 37 °C with 5% CO₂. Cells were infected with DENV2 at a multiplicity of infection of 1 for 1 h, after which the virus inocula were then removed and replaced with 5% FBS/DMEM maintenance media. At 6 h post-infection, the infected cells were treated with 7.5 μM NS3 peptides complexed with 22.5 μM penetratin in a molar ratio of 1:3 or 7.5 μM NS3 penetratin fusion peptides (Table 1). At 24 h post-infection, the cells were washed once with PBS prior to lysis by TRIzol for cellular viral RNA quantification by real-time RT-PCR analysis with primers that binds to the NS5 gene (forward 5′-CCGCTGACATGAGTTTGTGAGTC-3′ and reverse 5′-CATGACAGGAGAATCAACCG-3′) (40).

**ATPase Activity Assay**—The assay was carried out as described (13), with slight modifications. Purified NS2B2(320–368)NS3 WT or NS50A protein of 2.5 nM was preincubated at 37 °C with poly(U) (10 μg/ml) in 40 μl of reaction buffer (50 mM Tris/HCl, pH 7.5, 2 mM MgCl₂, 1.5 mM dithiothreitol, 0.05% Tween 20, 0.25 μg/ml bovine serum albumin (Sigma)) for 5 min. The reaction was initiated by the addition of 10 μl of varying ATP concentrations (2-fold serial dilution, starting from 2000 μM) and carried out for 10 min at 37 °C. 10 μl of malachite green reagent (BioAssay Systems) was added to stop the reaction. Absorbance was read at 635 nm after 30 min at room temperature. The KD of the protein was determined with GraphPad Prism 5, with Michaelis-Menten Equation 1,
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\[ V_o = (V_{max}[S]) / (K_m + [S]) \]  

(Eq. 1)

SAXS—SAXS data of the NS3(172–618) and NS3(172–618)-NS5(320–341) were measured by the NanoStar™ instrument (Bruker), equipped with a METALJET™ x-ray source and Vantec 2000-detector system. The METALJET™ source uses the liquid gallium source to deliver a high intensity x-ray beam at the wavelength of \( \lambda = 1.34 \, \text{Å} \). The SAXS measurements were carried out with the source to sample distance of 145 cm, a two-pinhole collimation system, and the sample to detector distance of 67 cm (41). SAXS experiments of both proteins were carried out at 1.2, 2.2, and 4 mg/ml in a sample volume of 40 \( \mu l \) at 15 °C. For each sample, a total of nine measurements at 5-min intervals were recorded. The data were flood-field and spatially corrected, and processed using the built-in SAXS software. We tested the possible radiation damage by comparing all data sets, and no changes were detected. The scattering intensity of the buffer was subtracted, and the difference curves were scaled for the concentration. All the data processing steps were performed automatically using the program package PRIMUS (42). The forward scattering \( I(0) \) and the radius of gyration \( R_g \) were evaluated using the Guinier approximation (43). These parameters were also computed from the entire scattering patterns using the indirect transform package GNOM (44), which also provides the distance distribution function \( p(r) \). Ten low resolution models of NS3(172–618) or NS3(172–618)-NS5(320–341) were independently built by the program GASBOR (45). The spatial discrepancy (NSD), which is a measure of similarity between sets of three-dimensional points, was computed between all 10 reconstructions using the DAMAVER program (46). The reconstruction with the least NSD was selected for NS3(172–618) or the fusion protein NS3(172–618)-NS5(320–341). The ensemble optimization method (EOM) suite was used to select an ensemble of conformations that best fit the experimental data, and the dimensions of selected conformations were compared with the random pool to evaluate the flexibility and compactness of NS3(172–618)-NS5(320–341) (47, 48).

DENV2 Full-length cDNA Clone Construction and Site-directed Mutagenesis—To construct a full-length DENV2 cDNA clone (GenBank™ accession EU081177.1, cosmopolitan genotype; Fig. 4A), low passage virus stock was subjected to viral RNA extraction by RNeasy kit (Qiagen), and three cDNA fragments (fragment boundaries indicated by nucleotide numbers; Fig. 4A) covering the complete genome were amplified from viral RNA by RT-PCR using SuperScript III one-step RT-PCR kits (Invitrogen). Fragment 1 contained the SpHl restriction site, a T7 promoter sequence, and DENV2 cDNA nucleotides 1–4498, which also contained the KpnI restriction site. Fragment 2 spanned from the KpnI site (nucleotide position 4493) to the XbaI site (nucleotide position 6008). Fragment 3 spanned from XbaI site (nucleotide position 6003) to the 3' end of the genome (nucleotide position 10,723), containing the SacI site at the end of the genome. The PCR product of each cDNA fragment was digested and cloned into a pre-digested and modified low copy number plasmid pWSK29 (49). The plasmid was modified by the replacement of the BssHII site that was located before the T7 promoter with the SpHl site by site-directed mutagenesis of the plasmid with the following primer: forward 5' - GGCAGTGGACATCGTAATACGAC-3' and reverse 5' - GTCAATACGGATCTGCTGCGC-3'. The underlined nucleotide corresponds to the mutation that was being made. The subclone that maintained each fragment was named accordingly as follows: pWSK29 D fragment 1, fragment 2, and fragment 3, respectively, and each subclone was validated by DNA sequencing by 1st BASE DNA Sequencing Services (Singapore) before proceeding for assembly. Subsequently, fragment 2 was inserted into the subclone pWSK29 D2 fragment 1 at the KpnI and XbaI site to generate subclone pWSK29 D2 fragment 1 + 2. Finally, fragment 3 was inserted to generate subclone pWSK29 D2 fragment 1 + 2 at XbaI and SacI sites to generate the full-length cDNA clone, pWSK29 D2 full length. The E. coli XL-1 Blue chemically competent cell (Strategene) was used for construction and propagation of the cDNA clones. Standard cloning procedures were performed with the exception that the cDNA clones were propagated at 30 °C for at least 20 h. All restriction enzymes were purchased from New England Biolabs.

The genome-length cDNA clones with NS3:N570A and NS5:K330A mutations were constructed using the subclone pWSK29 D2 fragment 3. The mutations were generated using QuikChange II XL site-directed mutagenesis kit (Stratagene) and performed according to the manufacturer's protocol. The following primers were used for the generation of both mutants: NS3:N570A forward (5'-CTTTTATGGAGTCAAGAACGCCCAATCTTTGGGAAGAAATG-3') and reverse (5'-CATTTTCTTTCCAGAGTTGCGGCTTGACTCCATCAAAG-3'); NS5:K330A forward (5'-TGTTAGGGCTGTCA- AACAGCACCTTGGGATGTCATCCCC-3') and reverse (5' - GGGGATGACATCCCCAAAGGTGGTGTTAGCAGCTAAC- CAC-3'). The underlined nucleotides correspond to the mutation that was being made. Mutations were confirmed by automated DNA sequencing, and fragment 3 bearing the mutation was excised from the plasmid by XbaI and SacI and inserted into subclone pWSK29 D2 fragments 1 + 2 that were similarly cut with XbaI and SacI.

In Vitro Transcription, RNA Electroporation, Plaque Assay, Real Time RT-PCR (Reverse Transcription-PCR), Immunofluorescence Assay (IFA), and Western Blot—BHK-21 cells were trypsinized, washed twice with cold PBS, and resuspended in Opti-MEM (Invitrogen) at a cell density of 1 \( \times 10^7 \) cells/ml. 10 \( \mu g \) of \textit{in vitro} transcribed RNA with T7 mMESSAGE mMACHINE kit (Ambion) of DENV2 WT and mutants were mixed with 800 \( \mu l \) of cell suspension in a pre-chilled 0.4-cm cuvette and electroporated at settings of 850 V and 25 microfarads, 2 pulses with an interval of 3 s. Electroporated cells were allowed to recover at room temperature for 10 min before resuspending in complete RPMI 1640 medium for cell recovery. Cells (3 \( \times 10^6 \)) were then seeded into a 12-well plate and incubated at 37 °C in the presence of 5% CO\(_2\). Media were changed to 2% FBS maintenance media after 6 h post-transfection. Samples were harvested every 24 h post-transfection until 120 h. Supernatants were collected and clarified for titering of the infectious virus particle by standard plaque assay and extracellular viral RNA quantification by real time RT-PCR analysis (40). Cells were then washed once with PBS prior to lysis by TRIzol reagent (Invitrogen) or 1 \( \times \) SDS-
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RESULTS

NS3(566–618) Interacts with NS5 RdRP—Even though NS3(303–618) of the helicase domain (residues 172–618) showed similar reactivity (Fig. 1A) as previously shown by Y2H study to interact with NS5(320–368) (bNLS) (36) of RdRP domain (residues 273–900) (35, 36), the amino acid residues of NS3 helicase that are responsible for binding to NS5 RdRP have not been pinpointed. However, it was hypothesized that helicase subdomain III, NS3(482–618), contains the interaction site (13, 22). To test this, we first established that the binding of NS2B, NS3 to coated NS5 RdRP increased in a dose-dependent manner as described previously (33) and carried out the NS3-NS5 interaction assay in a competitive ELISA format, with an increasing amount of the following competing proteins, namely NS3(172–618), GST-NS3(482–618) (subdomain III, or GST-NS3(566–618) (Fig. 1, A and B). From these data, all three NS3 truncated proteins were found to be able to compete with NS2B, NS3 protein binding to coated NS5 RdRP (Fig. 1C). The apparent $K_d$ values of the NS3-NS5 interaction for NS3(172–618), GST-NS3(482–618), and -(566–618) were found to be comparable (5.33 ± 0.35, 10.18 ± 0.91, and 10.25 ± 1.14 μM (mean ± S.D., n = 3) respectively) and also similar to the $K_d$ values for NS3(172–618) as in previous reports (33, 37). Next, we also determined the IC$_{50}$ value for NS3(172–618), which was 6.07 ± 2.60 μM. The IC$_{50}$ values for GST-NS3(482–618) and GST-NS3(566–618) were not determined as we were unable to reach 100% inhibition due to limited availability of truncated protein. Protein expression constructs spanning other NS3 subdomain regions (NS3(172–482), -(307–619), and (172–566)) were also generated in this study, but due to protein stickiness, protein instability, or low protein yield, these proteins could not be purified sufficiently for the competition assay.

NS3 Asn-570 Is Critical for the Interaction—To map the NS3 sequence that interacts with NS5 more precisely, we proceeded to screen an array of overlapping 15-mer peptides (Mimotopes) (53, 54) that spanned subdomain III in the same competitive ELISA format (Fig. 2A), and we identified two peptides (NS3(566–580) and -(571–585)) that moderately (p value = 0.06 and 0.009, respectively) blocked NS3-NS5 interaction, thus narrowing down the interaction region to residues 566–585 of NS3. We next tested the synthetic peptide NS3(566–585) in ELISA and showed that it could also disrupt NS3-NS5 interaction in a dose-dependent manner with an IC$_{50}$ value of 128.8 ± 2.57 μM (Fig. 2B). At the same time, another peptide, NS3(86–100) with a similar charge as NS3(566–585), did not compete. This indicates NS3-NS5 interaction involves sequence-specific residues on the NS3 protein. Additionally, the same peptide also inhibited NS3-NS5 interaction in the AlphaScreen assay format where the interacting partners were synthesized in vitro using the wheat germ expression system (data not shown) (55).

Next, we also used the peptide in the viral inhibition assay to determine whether blocking the interaction site could reduce viral replication (Fig. 2C). Uptake of NS3(566–585) peptide into infected cells was facilitated by a well characterized cell-penetrating peptide, penetratin (56), which forms a nonconvan-
lent complex with the peptide (57). As shown in Fig. 2C, NS3(566–585) peptide could reduce viral replication by ~33% (p value = 0.0195, Fig. 2C, panel i). As a positive control, we also tested the NS5(320–341) peptide that was covalently linked to penetratin, which can also facilitate the uptake of the peptide into infected cells (58). NS5(320–341) peptide could also reduce viral replication by ~33% (p value = 0.0043, Fig. 2C, panel ii). This suggests that blocking interaction between NS3 and NS5 could be a potential therapeutic target.

Sequence alignment of NS3(566–585) of DENV1–4 and other flaviviruses (Fig. 2D) suggested that Asn-570 (highlighted in gray) is highly conserved within this region and may be critical for the NS3-NS5 interaction. To test this, we synthesized NS3(566–585)(N570A) peptide and carried out the same competitive ELISA. We found that the replacement of asparagine by alanine at position 570 of NS3 resulted in a null-peptide with respect to its ability to block the NS3-NS5 interaction (Fig. 2B).

Next, we expressed and purified the NS3:N570A full-length protein to measure its ATPase activity (59), and we found that it was comparable with the WT NS3 protein (Fig. 2E). The affinity of ATP for NS3 WT and NS3 N570A was similar (K_m = 185.8 ± 10.86 and 176.1 ± 8.73 μM, respectively). Both proteins also had similar turnover numbers (k_cat = 3.15 and 3.11 s⁻¹, respectively). From these results, we surmised that Asn-570 of NS3 helicase subdomain III appears to be critical for the NS3-NS5 interaction without affecting the in vitro ATPase activity.

NS5(320–341) of RdRP bNLS Binds to NS3 Helicase in Solution—To gain some insights on the NS3-NS5 interaction at the structural level, we constructed a synthetic fusion of NS3(172–618) with NS5(320–341) (within bNLS) connected by a flexible linker (G4SG4, NS3(172–618)-NS5(320–341); Fig. 3A) for SAXS study. NS5(320–341) was selected based on surface plasma resonance data for peptide binding and NS5 peptide-phage ELISA that supported the interaction of NS5(320–341) to NS3(172–618) (data not shown). Through modifications of the purification strategy, NS3(172–618) and NS3(172–618)-NS5(320–341) were purified to high purity and used in the SAXS experiments, which provide three-dimensional low resolution structures in solution (60) as described for single proteins (41, 61, 62), as well as multidomain complexes.

**FIGURE 1.** Competitive NS3-NS5 interaction ELISA with NS3-truncated proteins. A, schematic representation and diagram of the recombinant NS3 and NS5 proteins that were expressed in E. coli and used in B for competitive NS3-NS5 interaction ELISA. C, during incubation of NS2B18NS3 protein with coated NS5 RdRP, 2-fold serial dilution of either NS3 or GST protein (starting from 20 μM) was added in triplicate competition experiments. GST protein was included as negative control. Data are shown as the mean ± S.D. of triplicates from two independent experiments.
like ATPase (63, 64) and the NS3 protease-helicase complex of DENV (11). SAXS data of both proteins at three different concentrations were collected to yield the final composite scattering curves shown in Fig. 3, B and C, which indicate that both proteins are monodispersed in solution. Inspection of the Guinier plots at low angles revealed good data quality and no protein aggregation (Fig. 3, B and C, insets). NS3(172–618) has a radius of gyration ($R_g$) of 25.16 ± 0.6 Å and a maximum dimension ($D_{max}$) of 72.95 Å (Fig. 3D), whereas the NS3(172–618)-NS5(320–341) has an $R_g$ of 25.42 ± 0.6 Å and a $D_{max}$ of 75.64 Å (Fig. 3D). Comparison of the forward scattering of both proteins with the values obtained from a reference solution of ATPase (63, 64).
lysozyme (14.4 kDa) yielded a molecular mass of 44.4 kDa for NS3(172–618) and 47.7 kDa for NS3(172–618)-NS5(320–341), indicating that both proteins are monomeric at the concentrations used.

The low resolution solution structures of NS3(172–618) and NS3(172–618)-NS5(320–341) were restored \textit{ab initio} using the program GASBOR (45). The normalized spatial discrepancy (NSD) of 10 independent reconstructions of NS3(172–618) is 0.95 ± 0.02 Å, and the NSD of 10 reconstructions of NS3(172–618)-NS5(320–341) is 0.99 ± 0.03 Å. The reconstructions with the least NSD from both proteins were selected, and the obtained solution shapes for both proteins yielded a good fit to the experimental data in the entire scattering range. The corresponding fits for NS3(172–618) and NS3(172–618)-NS5(320–341) have discrepancies of D_{max} = 1.15 and 1.04, respectively. NS3(172–618) and NS3(172–618)-NS5(320–341) revealed globular shapes, with the dimensions of about 59.9 × 54.4 × 34 and 65.6 × 57.7 × 36.8 Å, respectively. When superimposed, the crystallographic structure of NS3(176–618) (Protein Data Bank code 2JLS) accommodated very well in the

**FIGURE 3. SAXS of NS3(172–618)-NS5(320–341) indicates NS3-NS5 interaction.** A, schematic representation of recombinant NS3 and NS3-NS5 fusion proteins that were expressed in \textit{E. coli} and used in SAXS study. SAXS scattering pattern (○) and its corresponding experimental fit (——) of NS3(172–618) (B) and NS3(172–618)-NS5(320–341) (C) at protein concentrations of 1.2 mg/ml (red), 2.1 mg/ml (olive), and 4 mg/ml (blue). The curves are displayed in logarithmic unit for clarity. The insets in B and C show the respective Guinier plot. D, distance distribution functions of NS3(172–618) (——) and NS3(172–618)-NS5(320–341) (——). E, superposition of the determined solution shape of NS3(172–618) with the crystallographic structure of NS3(172–618) (Protein Data Bank code 2JLS (12)). Subdomains I, II, and III are colored in yellow, orange, and red, respectively. F, superposition of the NS3(172–618) and (green) and NS3(172–618)-NS5(320–341) (cyan) solution shapes. The \( R_g \) of the NS3(172–618) and NS3(172–618)-NS5(320–341) is 25.2 and 25.4 Å, respectively. The arrows indicate the two protrusions, one at the bottom of the NS3(172–618)-NS5(320–341) solution shape, which leads to a more elongated shape in this protein, reflected by the increased \( D_{max} \) value (see Fig. 3D). This protrusion is in proximity to the Asn-570 (blue sticks) and NS3 peptide region, NS3(566–585) (black schematic) (inset in F). The second protrusion of the NS3(172–618)-NS5(320–341) may be caused by a conformational alteration due to NS5(320–341) interactions. G, EOM/\( R_g \) distribution of the selected ensemble (red line) contains a narrow peak at 24.6 Å, which is slightly smaller than the center \( R_g \) of the random pool (gray filled area), 24.8 Å, suggesting NS3(172–618)-NS5(320–341) is rigid and compact in solution, and peptide NS5(320–341) is bound to NS3(172–618).
solution form of the protein (Fig. 3E), indicating the high quality of the solution data. When the solution form of NS3(172–618) (green) and NS3(172–618)-NS5(320–341) (cyan) were superimposed (Fig. 3F), two protrusions were observed and are denoted by arrows in Fig. 3F. Protrusion 1 that is at the bottom of the NS3(172–618)-NS5(320–341) solution shape leads to a more elongated conformation in NS3(172–618)-NS5(320–341) than NS3(172–618). This is also reflected by the increased \( \text{D}_{\text{max}} \) in NS3(172–618)-NS5(320–341) (75.64 Å) when compared with the \( \text{D}_{\text{max}} \) of NS3(172–618) (72.95 Å) (Fig. 3D). The protrusion is in close proximity to residue Asn-570 (shown as blue sticks in the inset of Fig. 3F) and to the NS3 peptide region, NS3(566–585) that used in ELISA (shown as black schematic in the inset of Fig. 3F). The arrangement of NS3(566–585) enables an interaction with NS5(320–341) to occur.

Comparison of the NS3(172–618) and NS3(172–618)-NS5(320–341) low resolution structures also revealed a second protrusion of the NS3(172–618)-NS5(320–341) that may reflect a conformational alteration due to the NS3(172–618) and NS5(320–341) interaction. Taken together, these results appear to be consistent with the ELISA data and suggest a physical interaction between NS3(172–618) and NS5(320–341) in solution.

To eliminate the possibility that the peptide may be flexible in solution, the x-ray scattering data set of NS3(172–618)-NS5(320–341) had been further analyzed using the Ensemble Optimization Method (EOM) (47, 48) to assess the compactness and flexibility of NS3(172–618)-NS5(320–341) in solution. Based on the width and position of the selected ensemble peak relative to random pool in \( R_g \) distribution, the flexibility and compactness of the protein in solution can be determined. In the case of NS3(172–618)-NS5(320–341), the \( R_g \) distribution of the ensemble contains a narrow peak, indicating that NS3(172–618)-NS5(320–341) is rigid in solution. This peak is centered at 24.6 Å, which is slightly smaller than the center \( R_g \) of a random pool, 24.8 Å, suggesting that NS3(172–618)-NS5(320–341) is compact in solution. The EOM data demonstrate that peptide NS5(320–341) is bound to NS3(172–618).

**NS3:N570A Mutant Has Reduced Infectious Virus Production and Viral Protein Synthesis**—To study the impact of NS3:N570A mutation on NS3-NS5 interaction during the virus life cycle, we first generated a DENV2 WT cDNA clone by standard molecular cloning techniques. DENV2 of strain D2/SG/05K3295DK1/2005 of the cosmopolitan genotype (GenBank™ accession number EU081177.1) was isolated during a local dengue outbreak that occurred in 2005 as part of Early Dengue Infection and Outcome (EDEN) Study in Singapore, and it was picked as the template for construction of the DENV2 full-length cDNA clone because of the well documented history of the patients (38). The overall schematic representation of cloning strategy is shown in Fig. 4A. The cDNA clone was subdivided into three fragments based on unique restrictions that are present within the genome to facilitate the assembly of the full-length clone. To prevent unwanted rearrangement of the clone, both full-length cDNA and cDNA fragments were maintained in low copy plasmid pWSK29 and *E. coli* XL-1 Blue that were transformed with these plasmids grown at 30°C (49). The assembled full-length cDNA clone contained a T7 promoter at the 5’ end for in vitro transcription and a SacI site at the 3’ end for linearization of cDNA. The use of SacI site generated the in vitro RNA transcript, which contains an additional nonviral nucleotide at the 3’ end, instead of two additional nonviral nucleotides for DENV cDNA that used the XbaI site (65). Prior to linearization for in vitro transcription, the full-length cDNA was checked for possible rearrangement by EcoRI digestion, and the predicted fragments of 9609, 4015, 1284, and 1122 bp were observed on 0.6% agarose gel (data not shown), indicating no sign of recombination (65).

IFA of BHK-21 cells that were transected with in vitro transcribed genome-length RNA showed increasing numbers of cells expressing E protein from days 1 to 3 post-transfection, with ~50–60% of cells being E-positive on day 3 (Fig. 4B). Plaque assay for supernatant showed an increase in infectious viral particles from days 1 to 3, peaking at 1 × 10^5 plaque-forming units/ml on day 3 (Fig. 4C). The size of the plaque on BHK-21 cells, which was produced by the supernatant of BHK-21-transfected cells, was comparable with those produced by the supernatant of C6/36-transfected cells (Fig. 4C, left and right insets, respectively). These results show that the RNA transcribed from the cDNA clone is highly infectious and that infectious virus can be produced in both BHK-21 mammalian cells and C6/36 mosquito cells.

After determining the growth kinetics of cDNA clone-derived DENV2 WT virus, the DENV2 NS3:N570A cDNA mutant clone was generated by site-directed mutagenesis, and its phenotype in infectious virus production and viral protein synthesis was examined over the course of 5 days’ post-transfection. For comparison, we also generated the NS5:K330A mutant (37), a known NS3-NS5 interaction-defective mutant that failed to replicate by reverse genetics studies, as a control. Similar to the NS5:K330A mutant, no infectious virus was recovered from the NS3:N570A mutant even when neat supernatant was used for titering (Fig. 5B). In agreement with the lack of infectious virus, extracellular viral RNA levels for both mutants were at the limit of detection of DENV RNA by RT-PCR and did not change over 5 days, suggesting that the mutants are inviable (Fig. 5A). The RNA detection for WT increased over 72 h when it reached maximal virus RNA detection that stabilized until 120 h. Next, to determine whether polyprotein synthesis from the transfected RNA is occurring and that the absence of infectious virus particles may be due to a lack of packaging and/or release of infectious virion, we checked the level of NS3 protein by both Western blot and IFA. Surprisingly, for the cells transected with NS3:N570A mutant transcript, we could detect NS3 protein using humanized monoclonal antibody 3F8 (33) in the 24- and 48-h samples (Fig. 5C, red arrows) but not in the 72-h sample. The highest percentage of NS3-positive cells (5–10%) was observed on day 3 post-transfection, which declined to ≤5% of NS3-positive cells on day 3 post-transfection (Fig. 5D), and finally an undetectable level at later time points on days 4 and 5 (data not shown). On the other hand, for the cells transfected with NS5:K330A mutant, the NS3 protein was almost undetectable by Western blot and IFA (<1% of NS3-positive cells) at all time points post-transfection (Fig. 5, C and D). Taken together, the results indicate that reduced NS3-
NS5 interaction in NS3:N570A mutant impairs infectious virus production and viral protein synthesis.

**NS3:N570A Mutant Showed Accumulation of Negative-strand RNA**—The reduction in viral protein synthesis for the NS3:N570A mutant and the lack of detectable infectious plaques suggest that attenuated RNA replication could occur at an early time point for this mutant. Therefore, we examined the RNA replication kinetics by real-time RT-PCR and IFA for detection of dsRNA. The results showed that intracellular RNA can be detected for NS3:N570A and WT but, intriguingly, not for NS5:K330A-transfected cells (Fig. 6A and B). The RNA copy for WT increased from around 10^6 copies/μg at 6 h post-transfection to 10^9 copies/μg of RNA at 120 h, as detected by real-time RT-PCR. Although less than WT, a 10-fold increase in intracellular viral RNA copy was observed from day 1 to 2 (p value = 0.008) for NS3:N570A mutant and thereafter, the level declined till day 5. The RNA copy for the NS5:K330A mutant declined from 6 to 48 h post-transfection and remained relatively stable from 48 to 120 h, suggesting that there was no viral RNA replication, and only the input RNA that was transfected by electroporation was being detected. Interestingly, the detection of the peak intracellular RNA level for NS3:N570A mutant on day 2 correlated with the highest level of NS3 protein detected by Western blotting (Fig. 5C).
To explore this further, we quantified the level of intracellular positive- and negative-strand viral RNA with strand-specific tagged primers in the real time RT-PCR experiment (Fig. 6C). Essentially, the tag sequence that was described by Plaskon et al. (51) was fused to the sequence that binds to E gene (50) to create a primer that can bind to either the positive or negative strand during the cDNA synthesis step. During real time RT-PCR, the tag and E-specific primers were used to distinguish cDNAs that were transcribed from either the positive- or negative-strand template for accurate quantification of both strands. As expected, for WT viral RNA-transfected cells, the level of both positive- and negative-strand RNAs increased over time until day 3 and declined at slightly different rates after that until day 5. The quantification was consistent with previous data that an excess of positive-over negative-strand RNA can be detected in WT virus-infected cells (66). The RNA quantification for the NS3:N570A mutant revealed a different trend to WT and NS5:K330A mutants. Overall, the level of positive- and negative-strand RNA synthesis for NS3:N570A was lower than WT. However, the level of negative-strand RNA synthesis (rate = \( -2.17 \times 10^5 \pm 5.82 \times 10^4 \) positive-strand/h) during a 6–24-h period, and at a somewhat similar rate for both strands (rate = \( 3.55 \times 10^5 \pm 3.74 \times 10^4 \) negative-strand/h and \( 7.48 \times 10^5 \pm 1.28 \times 10^5 \) negative-strand/h) during the 24–48-h period, suggesting that the negative-strand RNA may be accumulating as a double-stranded replicative form by base-pairing with the transfected positive-sense RNA. Interestingly, beyond 48 h, the rates at which both positive- and negative-strand RNA accumulated over time were almost identical. For NS5:K330A mutant, the level of detected positive- and negative-strand RNA corresponds to the amount of transfected positive- and negative-strand RNA that degraded over time. (Note: T7 transcribed positive-sense RNA used for transfection contained \( \sim 0.1–0.3\% \) negative-sense RNA that could be detected by the primers that bind to E gene and is the basis for the gray cutoff line for the residual negative-strand from transfected RNA in Fig. 6C.) The NS3:N570A mutant probably results in either reduced or completely abolished NS3-NS5 interaction and does not support further RNA synthesis from the dsRNA template that is formed (67). This is consistent with the detection of highest percentage of dsRNA-positive cells on day 2 (5–10%) by intracellular RNA staining that does not accu-
mulate as seen in WT at the later time points (Fig. 6B). Interestingly, our detailed analysis suggests that NS5:K330A mutant cannot synthesize negative-strand RNA, although the mutation has been shown to have no effect on the in vitro enzymatic activities (37). These results suggest that the impairment of infectious virus production and viral protein synthesis in the NS3:N570A mutant may be due to the inability of the mutant NS3 to engage with NS5 and use the dsRNA template to produce more positive-strand genomic RNA that can be translated and processed.

**DISCUSSION**

*Flavivirus* NS3 and NS5 proteins are recognized as attractive targets for antiviral drug development because of their important functional roles in viral replication (68–71). Because of the potential interdependence of the two proteins in orchestrating viral genome replication in the RC, the interaction between NS3 and NS5 has been proposed as a promising new target (28, 29, 37, 71). It was previously demonstrated by the Y2H study that NS3(303–618) of the helicase domain interacts with NS5(320–368) of the RdRP domain. Subsequently, based on available crystal structures, subdomain III of NS3 helicase (residue 483–618) was suggested to be involved in the interaction with NS5 because it has a large protein surface area, located away from the region of main catalytic active sites of NS3 (13, 22, 36). The NS5 residue Lys-330, which is located on the surface of NS5 thumb subdomain, has been identified as a critical residue in interacting with NS3 helicase (37).

In this study, we focused our attention on identifying the NS3 residues involved in the NS3-NS5 interaction, and we used biochemical, genetic, and biophysical approaches to investigate the functional relevance of the proposed interaction site. Through the use of truncated NS3 protein constructs, overlapping peptides, and peptide phage display, we had fine-mapped the interaction region to residues 566–585 for NS3 and residues 320–341 for NS5. Our attempts to obtain crystals of NS3 and NS5 in complex were not successful, and therefore, we employed SAXS to obtain a solution shape that supports the physical interaction between NS3(172–618) and NS5(320–341). Through EOM analysis of the SAXS data, we were able to demonstrate that NS3(172–618)-NS5(320–341) is compact and rigid, indicating that peptide NS5(320–341) makes specific and discrete interactions with the helicase domain, NS3(172–618).

Electrostatic potential surface analysis of NS3(566–585) and NS5(320–341) shows a negatively charged contiguous surface...
on NS3 and a positively charged contiguous surface on NS5, which are charge-complementary to each other, and this further supports the results that the two regions can indeed interact with one another via charge interaction (Fig. 7A). Inspection of the amino acid sequence in NS3(566–585) region within flavivirus (Fig. 2D) indicated a high conservation of sequence, and we identified a conserved amino acid within this region, residue Asn-570, which we showed to be important for NS3-NS5 interaction; the mutation of this residue to alanine disrupts the in vitro NS3-NS5 interaction. When the same mutation was engineered into the DENV2 cDNA clone, it abolished infectious virus production, similar to NS5:K330A mutant (37). However, unlike the NS5:K330A mutant, the NS3:N570A mutant was able to synthesize low but unsustainable amounts of viral RNA and proteins. Comparison of the pattern of positive- and negative-strand RNA synthesis between WT, NS3:N570A, and NS5:K330A confirmed that the NS5:K330A mutant is completely inactive since transfected positive-strand RNA degraded over time, and the negative-strand RNA detected corresponded to background. The WT RNA-transfected cells showed synchronized synthesis of positive- and negative-strand RNA, which is in agreement with previous studies that have shown an excess of positive- to negative-strand RNA in DENV-infected cells (66). However, it is possible that the long term coordinated synthesis of positive- and negative-strand RNA requires a functional RC with optimum protein interaction affinities between NS3 and NS5 (Fig. 7B). This is supported by the NS3:N570A mutant that showed fairly robust negative-strand synthesis using the transfected RNA as a template, and it is probably analogous to the situation at the early stages of viral replication in an infected cell. The weakened or abolished interaction between NS3 and NS5 in the NS3:N570A mutant does not support the replication of new positive-strand RNA from 6 to 24 h when compared with WT virus, which fits rather well with the carefully conducted real time RT-PCR quantifications. Surprisingly, the NS5:K330A mutant that has been shown to be enzymatically active in vitro and demonstrated to have no interaction with NS3 did not show a

FIGURE 7. Schematic representation of NS3-NS5-RNA interaction. A, surface electrostatic potential presentation of NS5(273–900) (RdRP) and NS3(172–618) (helicase). The protein backbones of NS5(320–341) and NS3(566–585) are shown in ribbon presentation. The side chains of NS5 Lys-330 and NS3 Asn-570 are displayed. B, simplified schematic model of NS3-NS5 interaction complex with RNA. Dashed line (green) is used to denote that the uncertain path that exiting template RNA from NS3 takes to enter NS5 for complementary daughter strand synthesis. Dark blue line denotes the unwound parental strand; green line denotes the newly synthesized daughter strand.
similar increase in negative-strand RNA as compared with NS3: N570A mutant. There may be two possible reasons for the inactivity of the NS5:K330A mutant in making negative-strand RNA. First, it may be due to impaired intramolecular signaling between the two NS5 domains that probably coordinate the two functional activities of NS5 (72) required for in vivo polymerase activity. Second, its weak/abolished interaction with NS5 may affect the unwinding of secondary structures in the transected positive-sense RNA that is needed for negative-strand synthesis. Taken together, our study indicates that different NS3-NS5 interaction-defective mutants can impair infectious virus production, viral protein synthesis, and RNA replication to varying degrees, which is likely to be dependent on the importance of the amino acids that are involved in NS3-NS5 interaction, and also possibly the intramolecular interactions in NS5. It is interesting to note that the coordinated synthesis of positive- and negative-strand RNA at the early stages of replication can be further explored by studying NS3-NS5 interaction mutants displaying varying strengths/degrees of binding and also the contribution of intramolecular cross-talk between the domains of NS5 in strand-specific RNA synthesis in the RC (Fig. 7B). Importantly, we also note that although the NS3 residue Asn-570 is conserved among the four DENV serotypes and several members of flavivirus genus, it is not conserved in yellow fever virus NS3, which has a histidine in place of asparagine at this position in NS3 and tyrosine instead of lysine at position 330 of NS5 (for sequence alignment, refer to Figs. 1B and 5 in Ref. 36).

Overall, this study has identified a potentially new druggable target for the development of antiviral drugs to block NS3-NS5 interaction that is essential for viral replication. The available crystal structures of NS3 and NS5 together with in vitro assays for interaction can be used for in silico and high throughput screening campaigns to find lead molecules for antiviral drug development. Alternatively, because the NS3:N570A mutant genome can be translated to a low level and is able to synthesize negative-strand RNA, it may serve as a potential RNA-mediated vaccine, although the basis for this requires development of new technology platforms.

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