Role of RNA Interference (RNAi) in Dengue Virus Replication and Identification of NS4B as an RNAi Suppressor

Pavan Kumar Kakumani, a, b, c Sankeet Singh Ponia, a, b Rajgokul K. S, a, c Vikas Sood, a, b Mahendran Chinnappan, a Akhil C. Banerjea, a Guruprasad R. Medigeshi, a, b Pawan Malhotra, a Sunil K. Mukherjee, a Raj K. Bhatnagar a

International Centre for Genetic Engineering and Biotechnology, Aruna Asaf Ali Marg, New Delhi, India; National Institute of Immunology, Aruna Asaf Ali Marg, New Delhi, India; Translational Health Science and Technology Institute, Gurgaon, Haryana, India.

RNA interference (RNAi) is an important antiviral defense response in plants and invertebrates; however, evidences for its contribution to mammalian antiviral defense are few. In the present study, we demonstrate the anti-dengue virus role of RNAi in mammalian cells. Dengue virus infection of Huh 7 cells decreased the mRNA levels of host RNAi factors, namely, Dicer, Drosha, Ago1, and Ago2, and in corollary, silencing of these genes in virus-infected cells enhanced dengue virus replication. In addition, we observed downregulation of many known human microRNAs (miRNAs) in response to viral infection. Using reversion-of-silencing assays, we further showed that NS4B of all four dengue virus serotypes is a potent RNAi suppressor. We generated a series of deletion mutants and demonstrated that NS4B mediates RNAi suppression via its middle and C-terminal domains, namely, transmembrane domain 3 (TMD3) and TMD5. Importantly, the NS4B N-terminal region, including the signal sequence 2K, which has been implicated in interferon (IFN)-antagonistic properties, was not involved in mediating RNAi suppressor activity. Site-directed mutagenesis of conserved residues revealed that a Phe-to-Ala (F112A) mutation in the TMD3 region resulted in a significant reduction of the RNAi suppression activity. The green fluorescent protein (GFP)-small interfering RNA (siRNA) biogenesis of the GFP-silenced line was considerably reduced by wild-type NS4B, while the F112A mutant abrogated this reduction. These results were further confirmed by in vitro dicer assays. Together, our results suggest the involvement of miRNA/RNAi pathways in dengue virus establishment and that dengue virus NS4B protein plays an important role in the modulation of the host RNAi/miRNA pathway to favor dengue virus replication.

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sembly also occurs through direct or indirect interactions between VSRs and the protein components of RISC. Fny-CMV2b protein physically interacts with the PAZ domain and part of the PIWI domain of Ago1 (20). A few VSRs use the GW/GW AGO hook to inhibit siRNA/miRNA-loaded RISC activity and also target the amplification of antiviral silencing. The vast majority of mammalian viral proteins have not been assessed for potential RNA silencing suppressor (RSS) activities.

Dengue virus (DV) is the causative agent of dengue fever, the most prevalent arthropod-borne viral illness in humans, with 50 to 100 million individuals infected annually worldwide. The role of the host RNAi pathway in mosquitoes upon arboviral infection has been extensively studied (21). The four serotypes of DV identified so far (DV1 to DV4) have a single-stranded genomic RNA of positive polarity that serves as the mRNA for the translation of a large polyprotein that is co- and posttranslationally processed. Most, if not all, nonstructural (NS) proteins are involved in the replication of dengue virus RNA. NS5 is the RNA-dependent RNA polymerase, NS3 acts as the viral serine protease as well as the RNA helicase, and the glycoprotein NS1 probably plays a role at an early step of viral RNA replication. Little is known about the functions of the small hydrophobic proteins NS2A, NS4A, and NS4B. NS4B is the largest of the small hydrophobic NS proteins of dengue virus, consisting of 248 amino acids (aa). It has been shown to be part of the viral replication complex and is also implicated in DV pathogenicity (22, 23).

Relatively little is known about the role of host RNAi/miRNA pathways, including the role of viral suppressor proteins, in flavivirus replication in mammalian cells (24). In the present study, we showed that DV infection downregulates components of host RNAi/miRNA machinery affecting the biogenesis of many known human miRNAs. Depletion of some of the RNAi components by siRNA-mediated silencing leads to increases in DV replication. This indicates that the RNAi machinery is involved in controlling DV replication in mammalian cells, and one or more of the DV proteins may counteract this host defense. We identified NS4B as one of the DV proteins with RNAi suppression activity and showed that this activity is independent of the interferon (IFN) inhibition functions of NS4B. Furthermore, the NS4B protein failed to bind dsRNA but interfered with dicing activity. Together, our results provide evidence for the antiviral function of the host RNAi/miRNA machinery in dengue virus replication in mammalian cells and implicate NS4B as a VSR that might promote virus replication.

**MATERIALS AND METHODS**

**Cell lines and viruses.** HuH 7 and HEK293T cells were propagated in Dulbecco’s modified Eagle medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 2 mM l-glutamine, 100 units/ml penicillin G sodium, and 100 μg/ml streptomycin sulfate and nonessential amino acids at 37°C and 5% CO2.

DV2 (P23085 INDI-60) was obtained from the National Institute of Virology, Pune, India, and propagated in C6/36 mosquito cells. Infected cell culture supernatant was used throughout the study.

**Gene synthesis and generation of plasmid constructs.** To express NS4B in the Sf21 cell line, the ns4b gene was amplified by using primers ns4b forward (5′-GGA ATG GGA AAC GAG ATG GGT TTT CTA CTA-3′) and ns4b reverse (5′-GTC GAC TTA CCT TCT TCT GTT GGT TGT GGT-3′). Plasmid pTM1.4-DV-2K (1–249) GFP (a kind gift of Ralf Bartenschlager, University of Heidelberg, Heidelberg, Germany) served as a template for PCR amplification of 2K-NS4B using primers 2K-ns4b forward (5′-GAA ATG GGA TTA ACC CCC CCA GAT AAC CAA TTG-3′) and 2K-ns4b reverse (5′-GTC GAC TTA CCT TCT TCT GTT GGT TGT TGT-3′). Additionally, NS4B sequences from the remaining serotypes of dengue virus were amplified by using the primers listed in Table 1. The amplified products were ligated into the pIB-V5-His/TOPO vector (Invitrogen, Grand Island, NY, USA) under the control of a baculovirus early promoter, OpIE2, for better expression levels.

To express NS4B in the leaves of Nicotiana xanthii, the ns4b gene was amplified by using primers ns4b forward (5′-GGA ATG GGA AAC GAG ATG GGT TTT CTA CTA-3′) and ns4b reverse (5′-GTC GAC TTA CCT TCT TCT GTT GGT TGT TGT-3′) and cloned into the plant binary vector pBluescript under the control of the cauliflower mosaic virus (CaMV) 35S promoter.

To express NS4B proteins of all serotypes and 2K-NS4B in a mammalian cell line (HEK293T), DNAs corresponding to these genes were amplified and cloned into pIB-V5-His/TOPO. The cloned fragments were mobilized into the mammalian expression vector pCDNA 3.1+ (Invitrogen, Grand Island, NY, USA) under the control of the human cytomegalovirus (CMV) promoter.

**siRNA transfection and virus infection.** Silencer Select siRNAs for Dicer, Drosha, Ago1, and Ago2 were purchased from Ambion. The knockdown studies were performed by transfecting 10 nM siRNA duplex HuH 7 cells by using a reverse transfection protocol with Lipofectamine 2000 (Invitrogen, Grand Island, NY, USA). At 48 h posttransfection, cells were infected with virus at a multiplicity of infection (MOI) of 10. The virus was diluted in DMEM containing 2% FBS, and cells were incubated for 1 h at 37°C on a rocker. After 1 h, cells were washed twice with PBS and grown in complete growth medium. Cells were collected at 48 h postinfection by trypsinization, and the cell pellets were washed once with phosphate-buffered saline (PBS) and lysed in TRIzol (Invitrogen, Grand Island, NY, USA) for RNA isolation. RNA was prepared according to the manufacturer’s protocol. DNase I-treated total RNA was used for quantitative real-time PCR (qRT-PCR).

**Quantitative real-time PCR.** The comparative threshold cycle (Ct) method with SYBR green was conducted for mRNA quantification of host RNAi factors using primers described previously (25), and viral genomic RNA was quantitated by using TaqMan primer-probe mix as described previously (26). Genome copy numbers were normalized to glyceraldehyde-3-phosphate dehydrogenase (GAPDH) mRNA levels. Relative expression was calculated by using the comparative threshold cycle method. Similarly, miRNA quantifications were performed by using the TaqMan microRNA assay system according to the manufacturer’s instructions.

**Small RNA library preparation and sequencing.** HuH 7 cells were infected with DV2 at an MOI of 10. At 48 h postinfection, the cells were harvested, and total RNA was extracted by using TRIzol according to the manufacturer’s protocol. The total RNA of the uninfected and infected cells was used to prepare a small RNA library using True Seq small RNA library preparation by Illumina Inc. The PCR products of the library corresponding to a length of 145 to 160 bp were considered and sequenced by using Solexa, a massively parallel sequencing technology.

**Analysis of sequence reads for known miRNAs.** Small 36-nucleotide (nt) RNA reads were produced by using the Illumina genome analyzer. Low-quality reads were trimmed with our own perl script. Adaptor sequences were accurately clipped with the aid of a dynamic programming tool.
algorithm. After elimination of redundancy, sequences of at least 18 nt and <26 nt were considered for subsequent analysis. Sequences that perfectly matched the reported human miRNA sequences were considered for subsequent analysis.

**Generation and overexpression of NS4B mutants.** To generate NS4B deletion mutants, ns4b gene segments with transmembrane domain (TMD) deletions were amplified by using specific primers. The primers used for the generation of the deletion mutants are listed in Table 2. The amplified products were ligated into the pIB-V5/His-TOPO vector, and the orientation of the genes was confirmed by PCR and restriction analysis. Conserved and alanine scan mutants in the TMD3 region of the ns4b gene were constructed by using the QuikChange site-directed mutagenesis kit (Agilent Technologies, Santa Clara, CA, USA) according to the manufacturer’s protocol. Briefly, PCR was performed on the wild-type (wt) ns4b gene cloned into a pIB-V5/His-TOPO vector using primer pairs with the corresponding mutation. *Pfu* Turbo DNA polymerase (Agilent Technologies, Santa Clara, CA, USA) was used for PCR. The mismatched oligonucleotides used for site-directed mutagenesis are shown in Table 3. After amplification, the PCR mixture was subsequently treated with DpnI enzyme to remove methylated parental DNA prior to transformation into *Escherichia coli* XL1B cells. Three clones were selected for each mutant, and the plasmid from each colony was isolated and sequenced to confirm the incorporation of the desired mutation. All of the deletion and substitution mutants were further cloned into pCDNA3.1+ for the incorporation of the desired mutation. The GFP expression in the transfected lines was confirmed by FACS analysis.

**Substitution-of-RNAi assay in a mammalian (HEK293T) cell line.** The suppression of the reporter gene expression in mammalian cell lines was quantified by FACS analysis. For the FACS analysis, Sf21 cells or HEK293T cells were washed with FACS-grade PBS (BD Biosciences) and resuspended in 400 μl of PBS. The fluorescence of the cells was determined by FACSQuant flow cytometry (BD, Franklin Lakes, NJ, USA). All the experiments involving quantification of GFP-fluorescing cells by FACS analysis were carried out three times, and statistical significance of the results was determined by the use of Student’s *t* test.

**IFN assay.** For the IFN assay, HeLa cells were considered instead of HEK293T cells. Unlike HEK293T cells, HeLa cells express TLR3 (Toll-like receptor 3) on their surface, which can recognize poly(I·C) to stimulate an IFN response inside the cell. The cells were cotransfected with 100 ng of plasmids encoding wt NS4B, 2K-NS4B, and domain BC/NS4B. Subsequently, at 24 h posttransfection, cells were transfected with poly(I·C) (20 μg/ml) by using Lipofectamine 2000 CD reagent (Invitrogen, Grand Island, NY, USA). The cells were lysed 7 h after the poly(I·C) treatment, and the luciferase expression level was determined by using the dual-luciferase reporter assay (Promega, Madison, WI, USA) according to the manufacturer’s protocol.
Phosphorimager screen (Amersham Biosciences, USA) overnight and SDS and once in 6 H11003. 

Preparation of gfp dsRNA. For both in vitro RNA-binding and dicing assays, the substrate, gfp dsRNA, was prepared by using the Riboprobe Combination System–SP6/T7 RNA polymerase (Promega, Madison, WI, USA), but for the RNA-binding assay, the substrate was incorporated with [α-32P]UTP. At the end of the reaction, the substrate was subjected to RNase-free DNase I digestion and purified on an Illustra MicroSpin G-25 column (GE Healthcare, Piscataway, NJ, USA).

Electrophoretic mobility shift assay. Electrophoretic mobility shift assays (EMSAs) were performed at 37°C for 20 min with 25,000 cpm per reaction and 3 μg of poly(dI·dC) (Sigma-Aldrich, St. Louis, MO, USA) in 1× binding buffer. The binding buffer contained 20 mM HEPES (pH 7.5), 1 mM dithiothreitol, 1 mM EDTA (pH 8.0), 1.5 mM MgCl2, and 4% glycerol in a final volume of 30 μl. The RNA–protein complex was resolved on a 6% acrylamide nondenaturing gel ( prerun for 1 h at 4°C) in 1× Tris-borate-EDTA (TBE) buffer at 200 V at 4°C. Gels were dried and exposed for autoradiography. Scanning was performed by using the Typhoon 9210 variable-mode imager (Amersham Biosciences).

Northern blotting. Total low-molecular-weight RNA was isolated from the GFP-expressing, GFP-silenced, and GFP-reverted cells by using a mirVANA miRNA isolation kit (Ambion, Austin, TX, USA). Ten micrograms of each low-molecular-weight RNA was resolved by electrophoresis on a 20% polyacrylamide gel containing 8 M urea. RNA was electroblotted for 90 min at 15 V onto a Hybond-N + membrane (GE Healthcare, Piscataway, NJ, USA) and immobilized by UV cross-linking at 1,200 mW of light. The membrane was washed three times in 6× SSC (1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate) with 0.2% SDS and once in 6× SSC with 0.2% SDS at 42°C and then exposed to a phosphorimager screen (Amersham Biosciences, USA) overnight and scanned at 200 μm with a Typhoon-9210 instrument (Amersham Biosciences, USA).

In vitro Dicer assay. dsRNA of ~900 bp of the GFP gene was transcribed in vitro by using the Riboprobe Combination System–SP6/T7 RNA polymerase (Promega, Madison, WI, USA). dsRNA was digested into siRNA (~22 bp) by recombinant human Dicer (Genlantis) either in the absence or in the presence of decreasing amounts of recombinant protein NS4B, F112A (29). Reaction products were loaded onto an RNase-free 3% agarose gel and stained with ethidium bromide.

RESULTS

RNAi factors modulate dengue virus replication in a human cell line (Huh 7). Although host RNAi machinery has been shown to restrict dengue virus replication in insect cell lines (30, 31), limited data are available on the anti-dengue virus response of RNAi machinery in mammalian cells (24). To examine the role of host RNAi in dengue virus replication, the main catalytic components of host RNAi, namely, Dicer, Drosha, Ago1, and Ago2, were knocked down by the corresponding siRNAs in Huh 7 cells, and the levels of knockdown were analyzed by qRT-PCR of the corresponding mRNAs in transfected cells (Fig. 1Ai). We then carried out infection of the knocked-down cells with DV and consequently measured DV RNA from cells at 48 h postinfection. A 6-fold increase in DV genomic RNA levels was observed in the cases of Drosha, Ago1, and Ago2 knockdowns, and a nearly 2-fold increase was seen in the case of Dicer at 48 h postinfection (Fig. 1Aii), compared to cells transfected with siRNA corresponding to a nontarget control (NTC). We further assessed the effect of DV infection on the host RNAi pathway by measuring the mRNA levels of these components in DV-infected cells. As shown in Fig. 1B, we observed a >65% reduction in the levels of these mRNAs, suggesting a negative regulation of RNAi machinery by DV infection. These data support the notion that downregulation of RNAi factors helps dengue virus multiplication.

Similar results were observed in the case of West Nile virus (WNV) replication upon siRNA knockdown of RNAi components. As shown by our findings, there was an enhancement of WNV replication in the cases of Drosha and Ago1 knockdowns (Fig. 1B), although the levels were not high compared to levels of dengue virus replication. These results indicated that RNAi machinery modulates dengue virus replication in mammalian cells, and the pattern may be conserved among flaviviruses.

Host response to dengue virus infection. The fact that Dicer, Drosha, Ago1, and Ago2 knockdowns impacted viral replication in mammalian cells (Fig. 1) suggested that both the cellular RNAi and miRNA pathways might be influenced by viral infection in mammalian cells. Since miRNAs have been shown to play an important role(s) in cellular development, metabolism, and proliferation, it is likely that the profile of host miRNAs might alter to cope with viral pathogenesis (11). Therefore, we examined the human miRNA responses to dengue virus infection by examining the changes in the host miRNA profile in cells infected with DV at 48 h postinfection. The population of small RNAs extracted from both the uninfected and infected cells was sequenced separately by using the Illumina next generation sequencing (NGS) platform. The total number of reads was normalized to 1 million for each sample for analysis purposes. The NGS data analysis revealed that the abundance of a majority of known human microRNAs had changed following DV infection. Of the total (151) known microRNAs randomly examined, 143 (94.7%) were downregulated, 3 (1.9%) (hsa-miR-149-5p, hsa-mir-204-5p, and hsa-miR-375) were upregulated, and 5 did not change. These results are not unexpected in view of the general downregulation of drosha and dicer levels following DV infection. A few of the downregulated miRNAs are displayed in Fig. 2Ai to iv, and these results were once again confirmed by stem-loop qRT–PCR analysis for a few of these miRNAs, namely, hsa-miR-10b-5p, hsa-miR-21-5p, hsa-miR-375, hsa-miR-125a-5p, hsa-miR-148a-3p, and hsa-miR-375 (Fig. 2B). In addition, we performed stem-loop qRT analysis of some human miRNAs upon siRNA knockdown of RNAi components in Huh 7 cells. As seen from our results (Fig. 2C), similar reductions in miRNA levels were observed compared to changes in the miRNA profile upon dengue virus infection (Fig. 1B and 2C). These results clearly indicate that the host miRNA/siRNA pathway plays an important role in controlling viral replication in mammalian cell lines.

Dengue virus NS4B is a viral RNAi suppressor protein. Plant viruses have been shown to suppress the RNAi/miRNA pathway by producing virus-encoded RNAi suppressor proteins (32). A number of approaches that involve transient/stable expression of virus-encoded proteins in an RNAi sensor line expressing the reporter genes have been used to analyze viral suppressor proteins (33, 34). We have used similar approaches to test the RNAi sup-
pressor activity of a number of viral proteins (16, 27). In the present study, we analyzed the NS4B protein for its RNAi suppressor activity. NS4B was selected based on its previously demonstrated role in alpha/beta interferon signaling inhibition (22, 23, 35). To assess the RNAi suppressor function of NS4B, we used a previously established Sf21 RNAi sensor line (experimental line 1) that was used to screen RNAi suppression properties of FHVB2 (16). This cell line is particularly useful in discriminating between the interferon response and the RNAi pathway, because Sf21 cells lack the machinery to elicit an interferon response to viral infection or the presence of dsRNA/dsDNA-like structures in their cytoplasm. We transfected the Sf21 RNAi sensor line with NS4B/pIB at 48 h postinfection and analyzed the cells for GFP expression. As shown in Fig. 3Aii, NS4B showed a significant reversion in the expression levels of GFP, similarly to FHVB2, a known RNAi suppressor (experimental line 2 [GFP-reverted cells]). To further confirm the RNAi suppressor properties of NS4B, we cloned the ns4b gene into the plant binary vector pBI121 (Fig. 3Bi) and carried out reversal of silencing by agroinfiltration-mediated transient ectopic expression of NS4B in tobacco leaf tissues. The infiltrated leaves showed a good amount of reversal of GFP activity at 8 days postinfiltration, while the empty control vector failed to show such a reversal (Fig. 3Bii). The RNAi suppressor properties of NS4B were further confirmed in a GFP-expressing mammalian sensor cell line, HEK293T. The bicistronic vector Retro-Q, which encodes GFP and carries a short hairpin gfp RNA, was cotransfected with either a plasmid containing NS4B (pCDNA-NS4B) or the empty vector alone (Fig. 3Ci). Cotransfection of the NS4B expression plasmid dramatically restored the number of GFP-expressing cells, as was the case with pCDNA-FHVB2, a known RNAi suppressor-expressing plasmid. However, no such reversal was observed when only the empty control plasmid without any suppressor gene was used (Fig. 3Cii). Expression of NS4B in transfected HEK293T cells was confirmed by Western blot analysis (see Fig. 7v). Taken together, these results clearly demonstrated that the dengue virus NS4B protein acts as a suppressor of RNAi.
FIG 2 Host miRNA response to dengue virus infection. (A) Changes in the host miRNA profile were observed upon dengue virus infection at 48 h in Huh 7 cells. The abundance of downregulated miRNAs (>1.5-fold) are presented in terms of the normalized read counts (i to iv) in both uninfected and infected samples. (B) Validation by stem-loop qRT analysis of some of the miRNAs down- and upregulated upon dengue virus infection at 48 h (#, P < 0.006). (C) Stem-loop qRT analysis of host miRNAs that are affected upon siRNA knockdowns in Huh 7 cells (#, P < 0.005). Data shown in panels B and C are means ± standard deviations from three independent experiments. The # symbols indicate a statistically significant difference in terms of the P value.
Four serotypes of DV (DV1 to DV4) have been identified so far, and we compared the sequences of NS4B proteins of these serotypes. As shown in Fig. 3Di, differences were observed among the different serotypes with respect to the NS4B protein, especially at the N and C termini. Therefore, we tested the suppressor activities of all four NS4B proteins in a reversal-of-silencing assay using the HEK293T cell line. Interestingly, NS4B proteins of all serotypes showed similar RNAi suppressor activities (Fig. 3Dii), thus advocating a conserved role of the protein as a suppressor in the viral life cycle.

**NS4B deletion mutants reveal that transmembrane domains (TMD3 and TMD5) are responsible for RNAi suppression.** NS4B consists of 248 amino acids and contains a 23-residue signal sequence, 2K, at its N terminus that translocates NS4B into the lumen of the ER (2K-NS4B). The domains of NS4B and 2K-NS4B are schematically shown in Fig. 4A. We compared the RNAi suppressor activities of NS4B and 2K-NS4B in the HEK293T sensor line. As shown in Fig. 4B, both these proteins were able to reverse the effect of RNAi on the sensor lines of HEK293T cells effectively, thereby suggesting that the 2K signal sequence does not significantly influence the RNAi suppressor activity of NS4B.

DV NS4B is an integral membrane protein, and several membrane topology models of NS4B predicted the presence of five TMDs (36). To delineate the NS4B regions involved in RNAi suppression, we generated several NS4B deletion mutants encompassing the entire length of the ns4b gene and analyzed the RNAi suppressor activity of each mutant in a reversal-of-RNAi assay using the HEK293T cell line. Figure 4C shows the schematics of the deletion mutants employed in the present study. As evident from Fig. 4D, a deletion mutant that retains only the TMD1 and TMD2 regions (domain A) had negligible RNAi suppressor activity, while deletion mutant B, consisting of the TMD3 and TMD4 regions, and deletion mutant C, consisting of the TMD5 region, showed RNAi suppressor activities similar to those of wild-type NS4B. Thus, the TMD1 and TMD2 regions play no role in RNAi suppression. This observation is noteworthy, as the N-terminal region (domain A) of NS4B was shown previously to inhibit alpha/beta interferon signaling (37). Deletion of the extreme C-terminal region (aa 191 to 248) of NS4B slightly affected its RNAi suppressor activity. Together, the results of the deletion mutant analysis showed that the TMD3 and TMD5 regions of NS4B are required for the RNAi suppressor activity, either as an individual unit or as a combinatorial group. Thus, the domain for the interferon response is distinct from that for the RNAi suppressor activity of NS4B.

The TMD3 region of NS4B is important for its RNAi suppressor activity. A series of substitution mutations in the ns4b gene had previously revealed that the TMD3 region is required for virus replication (38). To investigate the role of TMD3 in RNAi suppressor activity, we further generated six substitution mutants in the NS4B TMD3 region, namely, P104L/A, T108I/A, and F112L/A, and confirmed their sequences with automated gene sequencing. Figure 5A shows the location of the substitution mutants generated in the present study. RNAi suppression studies using the reversal-of-RNAi assay in a mammalian cell line were performed with the six mutant plasmids. As shown in Fig. 5B, a double-nucleotide modification (T340G and T341C) resulting in the amino acid change of F112A caused a significant reduction (~40%) of RNAi suppressor activity of NS4B, while a change of F112L did not (Fig. 5B). Expression of the F112A NS4B mutant was also confirmed by Western blot analysis (see Fig. 7v). The other four substitution mutations, P104L/A and T108I/A, did not change the RNAi suppression activities of NS4B significantly (Fig. 5B). The results from a previous study (38) and the present study together suggest that the F112A substitution in the TMD3 region affects RNAi suppressor activity as well as viral replication.

**NS4B harbors distinct domains for IFN-antagonistic and RNAi suppressor activities.** DV NS4B has been shown to inhibit the IFN signaling cascade (23), and intriguingly, most identified VSRs of mammalian viruses also have IFN- or protein kinase R (PKR)-antagonistic properties that may be linked to their RNAi suppressor function (39–41). To delineate the NS4B domains required for IFN-antagonistic activity, a HeLa cell line was transfected with a firefly luciferase expression plasmid under the control of the ISRE (interferon-stimulated response element) promoter pISRE-Luc (Agilent Technologies, Santa Clara, CA, USA), and Renilla luciferase was used as an internal control to normalize the transfection efficiency among all the samples (Fig. 6A). As a positive control, firefly luciferase expression was induced by poly(I-C), which produces a stimulatory IFN response. Cotransfection of a plasmid harboring ns4b along with pISRE-Luc did not result in IFN-antagonistic activity. However, transfection with 2K-NS4B resulted in a small but consistent reduction in the luciferase expression level (Fig. 6B), thereby suggesting that the N-terminal region of NS4B (2K region) is required for its IFN-antagonistic activity. As the "BC" domain of NS4B did not show a reduction of the luciferase expression level, the RNAi suppression domain of NS4B is thus clearly distinct from the IFN-antagonistic activity.

**NS4B does not bind dsRNA.** Many VSRs have been shown to cause RNAi suppression by binding to dsRNA/siRNA (14–16, 18, 40, 42). Although NS4B has been shown to dissociate NS3 from single-stranded RNA (ssRNA), there has been no report of its interaction with dsRNAs/siRNAs (43). To test the NS4B activity for dsRNA/siRNA binding, we expressed the protein in...
FIG 4 Analysis of deletion mutants for RNAi suppressor activity of NS4B. (A) Schematic representation of predicted transmembrane domains of NS4B with and without its signal peptide 2K. (B) Bar graph representation of the FACS results for NS4B and 2K-NS4B in HEK293T cells, represented on the x axis, with the percentage of cells expressing GFP represented on the y axis (#, P < 0.0004). (C) Schematic representation of deletion mutants of NS4B. (D) Bar graph representation of the FACS results for different deletion mutants of NS4B in HEK293T cells, represented on the x axis, with the percentage of cells expressing GFP represented on the y axis (*, P < 0.0004; #, P < 0.005). Data shown in panels B and D are means ± standard deviations from three independent experiments. The * and # symbols indicate statistically significant differences in terms of the P value.
a heterologous *E. coli* expression system. The recombinant protein was purified to apparent homogeneity and was tested for dsRNA/siRNA binding by an electrophoretic mobility shift assay (EMSA). However, we did not observe any significant binding of NS4B with labeled dsRNA (Fig. 7iii), but the recombinant dengue virus NS3 protein shifted the labeled dsRNA very well in the same assay, signifying the affinity of NS3 for dsRNA. These results suggest that NS4B suppressor activity might not involve dsRNA binding.

**NS4B inhibits Dicer processing of dsRNA into siRNA.** In order to explore other mechanism of RNAi suppression mediated by NS4B, we looked at the possibility of blockage of siRNA biogenesis by NS4B. Hence, we performed Northern analysis for gfp siRNA in GFP reversion assays with expression of either NS4B or its F112A mutant. As observed from our results, NS4B reduced the biogenesis of GFP-siRNA (Fig. 7i, lane 3) whereas the F112A mutant failed to significantly inhibit siRNA generation (Fig. 7i, lane 4). The level of expression of the mutant NS4B protein was similar to that of the wt NS4B protein (not shown). Hence, the *in vivo* reduction of siRNA biogenesis by NS4B seems clear.

Also, we investigated the *in vitro* processing of long dsRNA substrates by dicer in the presence of NS4B and its F112A mutant. An *in vitro* dicer assay was performed according to the manufacturer’s instructions (Genlantis, USA), in the presence of recombinant NS4B F112A proteins with increasing concentrations. Figure 7iv (lane 3) reveals that 1 unit of human dicer was able to process the dsRNA substrates into siRNAs (~22 bp). However, this processing was inhibited by NS4B in a dose-dependent manner (lanes 5 to 8). While 1 μg of NS4B protein blocked dicer processing almost completely (lane 4), about 3 μg of the F112A protein blocked processing by not more than 10%. These results clearly indicate that NS4B suppresses the RNAi pathway by inhibiting siRNA biogenesis.

**DISCUSSION**

Increasing evidence has emerged to suggest that RNAi pathways are the evolutionarily conserved defense mechanism against pathogenic viral infections (3). In early reports, plant RNAi was shown to act as a systemic antiviral defense against cytoplasmic RNA and nuclear DNA viruses (1). As in plants, the defense function of the RNAi pathway was also shown in *Caenorhabditis elegans* and *Drosophila melanogaster*, where mutations in the RNAi factor(s) showed increased susceptibility of these organisms to infection by viruses (44). The existence of analogous systems in vertebrates has been questioned simply because of the prevalence of interferon-based defense mechanisms (3). Recently, a number
of studies of mammalian cells have advocated the possible role of RNAi pathways against animal viruses (3, 9, 11). Mutations in RNAi factors or perturbations of RNAi pathways have been shown to increase vesicular stomatitis virus (VSV), influenza A virus, and human immunodeficiency virus type 1 (HIV-1) titers (3). In addition, HIV-1 and human T-cell leukemia virus type 1 (HTLV-1) infections in mammalian cells cause host cell miRNA profile changes (45), thus suggesting the role of RNAi/miRNA pathways in viral infections. Recently, a detailed analysis of small RNA repertoires in vertebrate and invertebrate systems using six mammalian RNA viruses unequivocally advocated an interplay between viral small RNAs (vsRNAs) and virus infection in vertebrate systems (9).

Given the importance of siRNAs/miRNAs in establishing viral infections, in the present study, we looked for the role(s) of host miRNA/RNAi factors, especially Dicer, Drosha, Ago1, and Ago2, in establishing dengue virus infections in mammalian cells. siRNA-mediated knockdown of these genes resulted in significant increases in replication of the dengue virus genome. We also observed reduced mRNA levels of these genes in dengue virus type 2-infected Huh 7 cells, and similar effects were observed in the case of WNV replication upon siRNA knockdowns. Together, these results suggested a role of the RNAi machinery in limiting dengue virus replication. Similar results were previously observed in the cases of equine encephalitis virus, respiratory syncytial virus, and influenza virus (46–48). Although this is the first report of a link between host miRNA/RNAi machineries and dengue virus infection in a mammalian system, a number of studies have reported the role of RNAi machinery in the replication of flaviviruses, especially dengue virus, in insect cells (30, 31, 37). Since the proteins Dicer, Drosha, Ago1, and Ago2 seem to be involved in modulating dengue virus replication in human cell lines, as evident from the present study, it suggested that both siRNA- as well as miRNA-mediated pathways may play an important role(s) in dengue virus replication in mammalian cells.

We next analyzed the changes in miRNA profiles in dengue virus serotype 2-infected Huh 7 cells. As observed in the cases of human retroviruses, hepatitis B virus (HBV), HCV, and herpes-
We observed a considerable reduction in the human miRNA profile in infected cells. As the drosha and dicer levels were downregulated following DV infection, the reduction in the levels of human miRNAs of the virus-infected cells was not very surprising. Together, these results suggest that both RNAi and miRNA pathways probably regulate dengue virus replication and pathogenesis in mammalian cells.

To overcome antiviral RNAi/miRNA responses, many plant, insect, and human viruses have been shown to encode suppressor of RNA silencing, also referred to as VSR (3, 5–7, 11). In this study, we analyzed dengue virus NS4B for its RNAi suppressor properties. NS4B was selected because it has been shown to inhibit alpha/beta interferon signaling, and a number of recent reports suggest a link between the IFN response and RNAi machinery in regulating viral replication in mammalian cells (3). Along with NS4B, we also examined NS4B with signal peptide 2K (2K-NS4B) for suppressor activity using reversal-of-silencing assays in three independent systems: the SF21 line, the HEK293T line, and plant leaves that were silenced for the expression of the reporter protein GFP. Such transient assays have been applied successfully to identify many plant and animal RNAi suppressors (16,27, 33, 34). Both the NS4B and 2K-NS4B proteins were effectively able to revert back GFP expression in GFP-silenced lines, thereby highlighting the role of NS4B as an RNAi suppressor in three different sensor lines. Most importantly, NS4B proteins from all four serotypes of dengue virus, serotypes 1 to 4, were able to reverse RNA silencing to a similar extent. It thus appears that NS4B-mediated characteristics are probably conserved across the eukaryotic kingdom.

A consensus in silico model for the topology of NS4B suggested that it contains five trans-membrane domains, designated TMD1, -2, -3, -4, and -5. Furthermore, expression studies using plasmids containing different NS4B predicted TMDs (pTMDs) fused to enhanced GFP (eGFP) at their COOH termini showed that the sequences containing pTMD3, -4, and -5, but not those contain-

![FIG 7](image-url)
ing pTMD1 and -2, mediate membrane targeting of eGFP (36). Based on these analyses, a number of NS4B deletion mutants were generated, and their suppression activities were analyzed. The NS4B deletion mutants retaining pTMD3 and -5 reversed RNA silencing to a level similar to that shown by wild-type NS4B. This observation is significant because the NS4B anti-IFN function is mediated by the pTMD1 and -2 regions of NS4B (35). Hence, NS4B showed the unique characteristic of possessing distinct domains for IFN antagonism and RNAi suppression.

Dengue virus NS4B has been shown to interact with the NS3 protein, and this interaction dissociates NS3 from single-stranded RNA. Furthermore, it has been shown that a single-amino-acid mutation in pTMD3 of NS4B (P104L), which had a pleiotropic effect on dengue virus replication in mosquito, disrupted this interaction (49). An independent study recently showed that two mutations in the NS4B pTMD3 region (C7152U and G7165U) attenuated the growth of wild-type DV1 and restored the replication of replication-defective dengue virus type 1 bearing a mutation in the C-terminal cytoplasmic portion of NS4A (38). These findings thus suggested that NS4B, particularly TMD3, contributes to the adaptation of DV for efficient replication. Viral replication is also linked with RNAi and its suppression. Thus, to investigate the details of the TMD3 involvement in RNAi suppression, we generated six different substitution mutants, P104L/A, T108I/A, and F112L/A, in the NS4B pTMD3 region and tested the suppressor activity of these mutant proteins. Three of these substitutions were the same as those described previously (38), while the other three were alanine substitutions for the same amino acids. Surprisingly, two substitution mutations, namely, P104L and F112L, which attenuated the growth of wild-type dengue virus serotype 1 in Vero cells in a previous study, did not have a substantial effect on the RNAi suppressor activity of NS4B. However, a substitution mutation, F112A, resulted in a considerable loss of RNAi suppression activity. These results are consistent with previous findings that suggested that NS4B, particularly its TMD3 region, contributes to the adaptation of DV for efficient replication in mammalian cells.

Experiments elaborating the mechanism of NS4B-mediated RNAi suppression revealed interesting insights into the mechanism of suppression. EMSA analysis with labeled dsRNA showed no affinity binding of NS4B. These results suggested that NS4B could be targeting one or a few components of the RNAi machinery for inactivation. The observed reduction in biogenesis of gfp siRNA from GFP reversion assays in HEK293T cells suggested the possible involvement of NS4B in dicing activity. This possibility was substantiated by the observed inhibition of dicer activity by NS4B in in vitro assays in a dose-dependent manner.

Furthermore, in vivo experiments revealed reduced dicer mRNA levels upon DV infection. In addition, our data suggest that dicer activity could also be reduced following viral infection. Thus, DVs have evolved many counterstrategies to defeat the RNAi-mediated host defense. It would be interesting to discover if any other RNAi components of the host are also affected by NS4B.

In summary, three major conclusions can be drawn from this work. RNAi/miRNA factors regulate dengue virus replication. Dengue virus NS4B acts as a viral RNAi suppressor, and the TMD3 region of NS4B, which has been shown to be involved in viral replication, is also important for its suppressor activity. We furthermore show that the NS4B protein suppresses the RNAi response by directly inhibiting the dicing process. As the profiles of human miRNAs change following virus infection, the abundance of RNAi factors might also consequently change, impacting virus replication and propagation indirectly. Thus, the modulation of DV replication within the human host could be due to the direct interference of NS4B in Dicer activity as well as the indirect effects mediated by the changes in the host microRNA profiles. Although the present study does not elaborate on the mechanism of NS4B suppression in detail, it is clear that NS4B suppresses RNAi by interfering with dicer activity. It is possible that NS4B might interact with another host RNAi factor(s) and inhibit the biochemical activities of the factor, resulting in an overall downregulation of the host RNAi response. Nevertheless, the present study provides a lead to elucidate such mechanisms and also advocates the development of inhibitors against NS4B so that viral replication can be blocked by chemical means.

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