Long-Range RNA-RNA Interactions Circularize the Dengue Virus Genome

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Secondary and tertiary RNA structures present in viral RNA genomes play essential regulatory roles during translation, RNA replication, and assembly of new viral particles. In the case of flaviviruses, RNA-RNA interactions between the 5′ and 3′ ends of the genome have been proposed to be required for RNA replication. We found that two RNA elements present at the ends of the dengue virus genome interact in vitro with high affinity. Visualization of individual molecules by atomic force microscopy revealed that physical interaction between these RNA elements results in cyclization of the viral RNA. Using RNA binding assays, we found that the putative cyclization sequences, known as 5′ and 3′ CS, present in all mosquito-borne flaviviruses, were necessary but not sufficient for RNA-RNA interaction. Additional sequences present at the 5′ and 3′ untranslated regions of the viral RNA were also required for RNA-RNA complex formation. We named these sequences 5′ and 3′ UAR (upstream AUG region). In order to investigate the functional role of 5′-3′ UAR complementarity, these sequences were mutated either separately, to destroy base pairing, or simultaneously, to restore complementarity in the context of full-length dengue virus RNA. Nonviable viruses were recovered after transfection of dengue virus RNA carrying mutations either at the 5′ or 3′ UAR, while the RNA containing the compensatory mutations was able to replicate. Since sequence complementarity between the ends of the genome is required for dengue virus viability, we propose that cyclization of the RNA is a required conformation for viral replication.

Outbreaks and epidemics caused by dengue virus continue to pose a public health problem in tropical and subtropical regions (60). It is estimated that more than 50 million human infections occur annually, and 2.5 billion people are at risk of dengue virus infection worldwide. Despite the wide morbidity and mortality associated with dengue virus infections, the molecular biology of this virus is not well understood, and at present, neither specific antiviral therapy nor licensed vaccine exists. Thus, defining the molecular determinants that regulate utilization of the viral RNA in the infected cell is of central importance for understanding the dengue virus life cycle.

The genomes of positive-strand RNA viruses participate in at least three different processes in the cytoplasm of the infected host cell: they serve as mRNA to direct the synthesis of viral proteins, they act as a template for genome amplification, and they are packaged along with structural proteins during viral assembly. The molecular mechanisms controlling the utilization of the viral RNA in each step of the viral life cycle are still poorly understood. Several lines of evidence support the notion that viral RNA genomes could circularize to regulate the initiation of translation and RNA synthesis at the 5′ and 3′ ends of the genome (4, 15, 18, 22, 23, 30, 31, 33, 38, 44). However, the molecular nature of 5′-3′ associations and the details of how different conformations of the RNA participate in the viral replication pathways remain unclear.

Different strategies for 5′-3′-end contact in the genome of positive-strand RNA viruses were proposed to be mediated by RNA binding proteins. Viral RNAs bearing a 5′-cap structure (m7GpppN) and a 3′ poly(A) tail could circularize by a protein bridge between eIF4G and eIF4E from the cap-binding complex and the poly(A) binding protein (PABP) similar to that observed for cellular mRNAs (59). Other strategies were proposed for viruses lacking the cap structure and/or the poly(A) tail. For instance, picornaviruses do not have a cap at the 5′ end but instead use a highly structured 5′-untranslated region (UTR) to bind the cellular poly(C) binding protein which mediates the interaction with PABP and the viral poly(A) tail (18, 30, 56). The viral protein NSP3 is a candidate to mediate interactions between the ends of the rotavirus genome. This protein binds the 3′ end of the viral RNA and the cellular protein IF4G of the cap-binding complex (46, 54). In the case of viral bovine diarrhea virus, which lacks both the cap structure and the poly(A) tail, binding of NFAR proteins to the viral 5′ and 3′ UTRs was found to mediate contacts between the 5′ and 3′ ends of the RNA (33). A different strategy was postulated for circularization of flavivirus genomes. In this case, direct RNA-RNA interactions between sequences present at the 5′ and 3′ ends of the viral genome have been proposed (23, 37, 38, 62). For all-mosquito borne flaviviruses, within the 3′ UTR, there is a putative cyclization sequence (CS) (3′ CS), which is complementary to a sequence located in the capsid coding region near the 5′ end of the genome (5′ CS) (11, 23, 38, 51).

The genus flavivirus of the Flaviviridae family includes important human pathogens such as dengue virus, yellow fever
virus, West Nile virus, and Japanese encephalitis virus. As with other positive-strand RNA viruses, replication of flaviviruses proceeds along a two-step pathway in the host cell. After synthesis and maturation, the nonstructural proteins and the viral RNA form a replication complex that catalyzes the synthesis of the negative-strand RNA, which in turn is used as a template to amplify new strands of genomic RNA. Even though contacts between the ends of the genomes of positive-strand RNA viruses have not been directly demonstrated, for flaviviruses, there is functional evidence supporting the hypothesis that genome circularization is required for viral replication (38).

The single-stranded RNA genome of flaviviruses is about 11 kb long and encodes one open reading frame flanked by 5′ and 3′ UTRs of about 100 and 600 nucleotides, respectively (49). The highly structured 3′ UTR ends in a very conserved 3′ stem-loop (SL), which is absolutely required for viral replication (8, 42, 47, 48, 64). The structure of the 3′ SL is preceded by the putative cyclization sequence 3′ CS. There is a total of 11 or 12 contiguous base pairings possible between 5′ and 3′ CS of West Nile, dengue, Japanese encephalitis, and Murray Valley virus RNAs (for a review, see reference 41) and 18 nucleotides in the case of yellow fever RNA (11). Computer analysis using the complete RNA of Kunjin virus indicates that base pairing of 5′-3′ CS is thermodynamically feasible (37). Using in vitro assays, it has been demonstrated that efficient RNA synthesis by dengue virus RNA-dependent RNA polymerase requires both the 5′ and 3′ CS (62). Interestingly, Khromyk et al. examined the importance of 5′ and 3′ CS complementarity using mutated Kunjin virus replicas (38). Specific mutations in 5′ or 3′ CS abolished RNA amplification, while reconstitution of the base pairing with foreign sequences restored viral replication, suggesting a functional role for 5′-3′-end contact in the viral genome.

In this article, we show that the dengue virus genome circu-
larizes through RNA-RNA interactions in the absence of pro-
tiens. Using atomic force microscopy (AFM), we visualized individual dengue virus RNA molecules in circular conforma-
tions. We found that sequences encompassing the previously reported 5′ and 3′ CS regions are essential but not sufficient for RNA interactions. We identified a new element of 16 nucleotides present at the 5′ and 3′ UTRs as an important determinant for RNA-RNA association. Importantly, func-
tional studies obtained here with recombinant dengue viruses strongly suggest that RNA complementarity between the ends of the genome is necessary for viral viability.

MATERIALS AND METHODS

RNA preparation. RNAs were obtained by in vitro transcription using T7 RNA polymerase (90 min, 37°C) and treated with DNase 1 RNase-free to remove templates. The RNAs were purified using an RNaseasy Mini kit (QIAGEN Inc.) to remove free nucleotides and quantified spectrophotometrically, and their integrity was verified by electrophoresis on agarose gels. All numbers given below in parentheses refer to nucleotide positions of a dengue virus type 2 strain 16681 infectious cDNA clone (GenBank accession number U87411).

The sequences corresponding to the 5′ UTR-C62 (nucleotides [nt] 1 to 160) and 3′ SL (nt 10617 to 10723) were amplified by PCR from a dengue virus infectious clone (40) with a forward primer carrying the T7 RNA polymerase promoter. Mutations in 5′ CS (mutations 143, 144, 145, and 146) were introduced by PCR using antisense primers carrying the desired mutation. Mutations in the 5′ upstream AUG region (UAR) (mutations 131, 133, 139, 170, and 175) and mutant 3′ SL 177 were generated by overlapping PCR. PCR products were directly used as templates for in vitro transcription. The position and the specific nucleotide substitution are indicated in Fig. 4, 6, and 7, respectively.

To generate the model RNA molecule of 2.3 kb, pGL5′3′D/5′UTR-C62 was constructed by introducing the 5′ UTR-C62 (nt 1 to 160) into pGL3-Basic vector (Promega) between SacI and Neot restriction sites and dengue virus 3′ UTR (nt 10629 to 10723) between XbaI and BamHI sites. The construct carrying dengue virus 5′ UTR-C62 and 3′ UTR-C62 was linearized using the enzyme SmaI, and its product was used as a template for in vitro transcription. Deletions of 96 nucleotides (nt 10627 to 10723, deleting the 3′ UAR) and 106 nucleotides (nt 10617 to 10723, deleting both the 3′ CS and UAR) at the 3′ end were introduced into the construct by PCR using pGL5′3′D as a template. The DNA template for antisense RNA synthesis was generated by PCR amplification of 1.6-kb or 1.8-kb fragments of luciferase sequence. Full-length dengue virus RNA was generated using as a template dengue virus type 2 strain 16681 infectious cDNA clone (pD2/IC-30P-A) linearized by the XbaI restriction enzyme. A DNA template to synthesize an antisense RNA molecule spanning NS4B and NS5 coding sequence (nt 6970 to 10721) was also generated by PCR.

RNA binding assays. RNA-RNA interactions were analyzed by electrophoretic mobility shift assays. Uniformly 32P-labeled RNA probes were obtained by in vitro transcription using T7 RNA polymerase and purified on 5% poly-
acrylamide gels and 6 M urea. The binding reaction mixtures contained 5 nm HEPES, pH 7.9, 100 mM KCl, 5 mM MgCl₂, 3.8% glycerol, 2.5 μg RNA, 5′ UTR-C62 (concentration indicated in each case), and uniformly 32P-labeled 3′ UTR-C62 RNA (0.1 μM, 30,000 cpm), in a final volume of 30 μl. RNA samples were heated at 85°C for 5 min and slow cooled to room temperature. RNA-RNA complexes were analyzed by electrophoresis through native 5% poly-
acrylamide gels supplemented with 5% glycerol. Gels were prerun for 30 min at 4°C and then loaded and then 25 μl of gel was allowed to proceed for 4 h at constant voltage. Gels were dried and visualized by auto-
radiography or exposed on a PhosphorImager plate.

To determine relative binding affinities, radiolabeled 3′ SL (0.1 M) was titrated with increasing concentrations of the unlabeled RNA as indicated in each case. The total radioactivity for each lane was determined by quantifying the radioactivity incorporated into the RNA-RNA complex plus the amount remain-
ing in the free probe. For each sample, we calculated the fraction bound, . The apparent Kₐ were estimated by fitting the data using nonlinear regression analysis as follows: = [RNA]([RNA] + Kₐ). Because the bound RNA was incor-
porated into two bands, we considered bound RNA as a single species equal to the sum of both bands.

AFM sample preparation and imaging. For AFM imaging, sense and antisense RNA molecules were denatured by heating at 85°C for 1 min and slow cooled to room temperature to form double-stranded RNA species in buffer A containing 20 mM HEPES, pH 8, and 4 mM MgCl₂. Samples were diluted to 1 ng/μl in buffer A and 20 μl of the mix was deposited onto freshly cleaved mica mica. After 2 to 5 min, the sample was gently washed with 0.5 ml mili0 water to remove molecules that were not firmly attached to the mica and blown dry with nitrogen. Tapping-mode AFM was performed using a NanoScope III Multimode (. Digital Instruments, Santa Barbara, CA) with the cantilever (T Apparent Kₐ were estimated to range from nanometers to 100 μm. Nanomaterials collagen microscleres 125 μm in length and a force constant of ~40 N m⁻¹ were used (NanoDevices, Veeco Metrology, Santa Barbara, CA). Cantilever oscillation frequency was tuned to the resonance frequency of the cantilever (280 to 350 kHz). After a period of 15 to 30 min of thermal relaxation, initial engagement of the tip was achieved at scan size zero to minimize sample deformation and tip contamination. The images (512 by 512 pixels) were captured with a scan size of 0.5 μm at a scan rate of 1 to 2 scan lines per s. Images were processed by flattening using Nanoscope software (Digital Instruments) to re-
move background slope. Measurements were done using Nanoscope software and ImageJ, version 1.3 (NIH).

Construction of recombinant dengue viruses. The full-length cDNA of dengue virus type 2 pD2IC/30P-A was modified by site-directed mutagenesis to generate a 3′ UTR cassette between unique AflII and XbaI restriction sites. To this end, PCR product generated with sense primer AVG-62 (5′-ACAACGTGTGGAGGA CATAGCATTGA-3′) and antisense primer AVG-91′ (5′-TATAGGAGCAAACTTAAGATGAAAC-3′) was fused by overlapping PCR. This PCR product was cloned into pGEM-T Easy (Promega), generating pGEM-3′UTR-C62. The AvrII-ClaI fragment of pD2IC/ 30P-A was replaced with the AvrII-ClaI fragment of pGEM-3′UTR4/3 to generate pD2IC/30P. The mutant DV-3′UTR177 was obtained by exchanging the wild-type fragment of AflII-XbaI with the XbaI restriction sites. To this end, DNA samples were generated using as a template dengue virus type 2 strain 16681 infectious cDNA clone (pD2/IC-30P-A) linearized by the XbaI restriction enzyme. A DNA template to synthesize an antisense RNA molecule spanning NS4B and NS5 coding sequence (nt 6970 to 10721) was also generated by PCR.

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(nucleotide substitutions are underlined). The mutant DV-5’UAR 175, containing the mutation 80-ACAAGACACAUCUGUG, was obtained by overlapping PCR using a 1,508-nucleotide-long fragment from the 5’ end of pD2/IC89. The PCR fragment was digested with Sacl and SphI, and the resulting fragment was cloned into the homologous restriction sites within pD2/IC89 or the DV-3’UAR 177 mutant clone to obtain DV-5’UAR 175 and DV-5’UAR 175-177 dRNAs, respectively.

RNA transfection and recovery of viruses. Wild-type (WT) or recombinant plasmid DNAs were linearized with XbaI and used as templates for transcription by T7 RNA polymerase in the presence of m7GpppA cap structure analog. RNA transcripts (3 μg) were transfected with Lipofectamine 2000 (Invitrogen) into BHK-21 cells grown in 60-mm-diameter tissue culture dishes. The transfected cells were trypsinized on day 3 posttransfection and two-thirds of the total cells were reseded. This procedure was repeated every 3 days for 21 days. Supernatants derived from transfected BHK-21 cells were harvested at 3, 6, 9, 12, 15, 18, and 21 days posttransfection and used to search for infectious dengue virus. For plaque assays, 3.0 × 10⁴  to 4.0 × 10⁵ BHK-21 cells were seeded per well in 24-well plates and allowed to attach overnight. Transfected cell supernatants were serially diluted, and 0.1 ml of the inoculum was incubated on the cells for 60 min. Afterwards, 1 ml of overlay (1% minimal essential alpha medium, 2% NCS, 100 U of penicillin/ml, 100 μg of streptomycin/ml, and 0.8% methyl cel- lulose) was added to each well. Cells were fixed 7 to 9 days postinfection with 10% formaldehyde and stained with crystal violet.

Immunofluorescence. Transfected cells with wild-type and mutated full-length dengue virus RNA were used for immunofluorescence assay (IFA). BHK-21 cells were grown in 60-mm-diameter tissue culture dishes containing a 1-cm² coverslip inside. The coverslips were removed and directly used for IFA analysis. The transfected cells were trypsinized at day 3, and two-thirds of the total cells were reseded to a 60-mm-diameter tissue culture dish containing a new coverslip inside. This procedure was repeated every 3 days for 21 days. At each time point, a 1:200 dilution of murine hyperimmune ascitic fluid against dengue virus type 2 in phosphate-buffered saline–0.2% gelatin was used to detect viral antigens. Cells were fixed in paraformaldehyde. Alexa Fluor 488 rabbit anti-mouse immunoglobulin G and Alexa Fluor 488 goat anti-rabbit immunoglobulin G conjugates (Molecular Probes) were used as detector antibodies at a 1:500 dilution. Photomicrographs (×200 magnification) were acquired with an Olympus BX60 microscope coupled to a CoolSnap-Pro digital camera (Media Cybernetics) and analyzed with Image-Pro Plus software.

Viral RNA extraction and sequencing. Viral RNA was TRIZOL extracted from a 200-μl aliquot of the media from transfected cells. Dengue virus RNA was reverse transcribed and amplified. Primers in the reverse transcriptase PCR (RT-PCR) were targeted to amplify nt 1 to 1451 in the 5’ end and nt 10201 to 10723 in the 3’ end. This primer design excluded the last 23 nt of the genome inside. The coverslips were removed and directly used for IFA analysis. The RT-PCR products were sequenced using an ABI 377 automated DNA sequencer and Big Dye terminator chemistry (Applied Biosystems).

RESULTS

Specific RNA-RNA interactions in the dengue virus genome. To investigate RNA-RNA association between different RNA elements of the dengue virus genome, we developed a method to examine the interaction of in vitro-generated RNA molecules using gel shift assays. For this purpose, radiolabeled RNAs corresponding to each of the four domains of dengue virus 3’ UTR (variable region, A2, A3, and 3’ SL) were in vitro transcribed. We evaluated the interaction of an RNA molecule corresponding to the 98-nucleotide-long 5’ UTR followed by 60 nucleotides of the coding region of the C protein (named here 5’UTR-C62 RNA) (Fig. 1A) with each of the four RNA probes. The RNAs were heat denatured, mixed, cooled to room temperature to allow folding, and used in gel shift assays. RNA-RNA complexes were observed only when the 5’UTR-C62 RNA was incubated with the RNA probe corresponding to the last 106 nucleotides of the viral genome (3’ SL RNA). RNA titrations indicate high affinity between the two RNA elements with an apparent dissociation constant (Kd) of 8 nM (Fig. 1B).

To determine specificity of the detected RNA-RNA complex, we measured the efficiency of complex formation in the presence of various amounts of iRNA or unrelated RNA (up to a 1,000-fold molar excess with respect to the unlabeled 5’ UTR-C62 RNA) at different ionic strengths (up to 500 mM KCl). Under these conditions, the RNA complex remained
unchanged, and it was chased only by specific unlabeled 3’ SL RNA (data not shown). Divalent cations are known to favor RNA tertiary structures under physiological ionic strengths (14, 43). Thus, we analyzed the effect of different concentrations of MgCl₂ and NiCl₂ on complex formation. We observed an absolute requirement of Mg²⁺ for RNA-RNA interaction (Fig. 1C). Mg²⁺ could not be replaced by Ni²⁺; in fact, in the presence of Ni²⁺, complex formation was reduced. In the presence of 100 mM KCl, 4 mM Mg²⁺ was required to allow 100% of the complex to be formed, suggesting that tertiary structures in the RNA are necessary for 5’ UTR-C62-3’ SL interaction. Taking these data together, by using in vitro RNA binding assays, it was possible to identify direct and specific RNA-RNA interactions between the 3’ SL and the 5’ UTR-C62 RNA.

**Atomic force microscopy reveals circularization of dengue virus RNA.** To examine whether the RNA-RNA interaction observed in gel shift assays was sufficient for mediating long-range interactions leading to single RNA molecule circularization, we used AFM (25). This technique allows visualization of individual molecules in physiological environments. In agreement with previous reports, single-stranded RNA molecules acquired compact conformations precluding visualization of intramolecular interactions (27, 36). In order to obtain RNA molecules suitable for AFM analysis, we used a method that

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FIG. 2. Single-molecule analysis reveals cyclization of an RNA molecule carrying the 5’- and 3’-end sequences of dengue virus. (A) Schematic representation of a model RNA molecule of 2.3 kb showing the 5’ and 3’ dengue virus sequences flanking the luciferase coding sequence. Annealing of an antisense RNA of 1,633 nucleotides is shown. The resulting molecule bears single-stranded overhangs in the 5’ and 3’ ends of 230 and 451 nucleotides, respectively. On the right, purified single-stranded RNA (ssRNA) and double-stranded RNA were resolved on a 1% agarose gel and visualized by ethidium bromide staining. (B) Visualization of the model RNA molecules by AFM. A single RNA molecule is shown in a linear conformation. The double-stranded RNA region is flanked by single-stranded regions corresponding to the 5’ UTR-C62 and 3’ UTR of dengue virus. (C) An image of a representative field of RNA molecules deposited on mica obtained by tapping-mode AFM. Circular, linear, and head-to-tail dimers were observed. (D) Image of individual RNA molecules in circular conformation is shown. Contacts between the 5’ and 3’ single-stranded regions of the molecules can be observed. (E) Schematic representation of the same RNA molecule shown in A hybridized with an antisense RNA molecule of 1 kb. The double-stranded region of 1 kb is flanked by a 5’ single-stranded region of 863 nucleotides that contained the 160 nucleotides of the 5’ end of dengue virus and a 3’ single-stranded region that corresponds to the 3’ UTR of dengue virus. At the bottom, a representative image of an individual molecule with a double-stranded region of 1 kb is shown in circular conformation.
was previously employed to visualize conformations of cellular mRNAs (59). This method consists of generating double-stranded RNA regions in the center of the molecule, leaving the sequences presumably involved in long-range interactions as single-stranded overhangs. Based on this report, we first developed a model RNA molecule of 2.3 kb carrying the 5′/H11032UTR-C62 and the complete 3′/H11032UTR of dengue virus flanking a luciferase coding sequence and a second RNA molecule of 1.6 kb complementary to the luciferase coding region. Hybridization of these two RNA molecules leaves the 5′ and 3′ dengue virus sequences of interest as single-stranded overhangs (Fig. 2A). The sense and antisense RNA molecules were in vitro synthesized, treated with DNases, purified, and mixed after thermal denaturation. Annealing of the two RNAs and integrity of the molecules were confirmed by agarose gels (Fig. 2A). The samples were deposited onto freshly cleaved mica and analyzed using tapping-mode AFM in air. High-resolution images of individual molecules were used to measure contour lengths of the double-stranded region as well as the apparent volumes of the 5′ and 3′ dengue virus sequences of interest as single-stranded overhangs. The contour length of the 1,633-nt double-stranded region was 437 nm with a rise per base pair of 0.27 nm. This observation is in good agreement with the values obtained for duplex RNA in the A form (50) and previous AFM data (7, 29). The single-stranded regions corresponding to the 5′ and 3′ UTR of dengue virus adopted globule-like conformations with apparent volumes of 25.7 nm³ and 131.4 nm³, respectively (Fig. 2B). Image analysis revealed the presence of both circular and linear conformations of the RNA. In Fig. 2C, we show a representative field (3 μm by 3 μm) containing molecules in different conformations, and Fig. 2D shows a higher-resolution image (0.7 μm by 0.7 μm) depicting circular conformations of the RNA. The molecules were categorized as being either circular or linear by visual inspection of a series of images from more than six independent experiments in which only intact molecules, determined by size, were counted. We estimated that 45% of the intact molecules were circular, and the remainder were linear (Table 1). At the concentration used for imaging (1 ng of RNA/μl), about 13% of the molecules were observed as RNA-RNA head-to-tail dimers. As a control, an RNA molecule with a deletion of the last 106 nucleotides (Δ3′SL RNA) was constructed, annealed with the 1.6-kb antisense RNA, and used for AFM imaging analysis. Visual inspection of this RNA revealed the absence of molecules in circular conformation (Table 1). Taken together, these observations indicate that long-range RNA-RNA contacts mediated by dengue virus sequences are capable of circularizing a model molecule. In addition, we confirmed that the last 106 nucleotides of the 3′ UTR are involved in RNA-RNA interaction, in agreement with the results obtained using RNA binding assays.

In these experiments, we were concerned about the possibility that the stiffness of the duplex RNA used in our model molecule could restrict RNA bending and circularization. Previous physicochemical studies performed with double-strand RNA polymers indicated that the persistence length (measure-

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* a Molecules were counted only when the apparent contour length of the double-stranded region was 437 ± 10 nm.

**FIG. 3.** Visualization of genome-length dengue virus RNA by AFM. (A) Schematic representation of the dengue virus genome. An antisense RNA anneals to the molecule, resulting in a 3,302-bp double-strand species with single-stranded regions of 6,970 and 451 nucleotides at the 5′ and 3′ ends, respectively. (B) Visualization of representative images of individual genome-length RNA molecules obtained by tapping-mode AFM.
ment of the flexibility of the molecules) was around 110 nm (35) and that the optimal loop size was three to four times the persistence length (330 to 440 nm) (34, 61). The double-stranded region used in our model molecule was 437 nm, which is around the optimal size to allow free movement of the ends of the molecule. Therefore, we conclude that the stiffness of the double-stranded region in the RNA molecules used in our experiments should not be a limitation for RNA circularization. To better characterize the system, we designed various antisense molecules that would leave single-stranded overhang regions of different lengths. All the molecules used (antisense from 1.6 to 1 kb) yielded similar results. Figure 2E shows the image of a representative molecule in a circular conformation with a double-stranded region of 1 kb.

Having shown that specific interactions between the 5' and 3' ends of dengue virus are capable of circularizing a model RNA molecule, we sought to analyze the conformation of the full-length dengue virus RNA. To this end, we in vitro synthesize...
For this reason, it is likely that the 25% of the molecules found in circular conformation was an underestimation. Nevertheless, the absence of circular conformations in the control RNA, lacking the 3' most 106 nucleotides, confirms that the amount of circular molecules observed with the genomic RNA is highly significant (data not shown).

Altogether, these results strongly support the hypothesis that specific dengue virus sequences mediate long-range RNA-RNA contacts in the viral genome.

Contacts between 5' and 3' CS are involved in RNA-RNA interaction. Nucleotide sequence comparisons between different flavivirus genomes indicate that the putative 5' and 3' CS are highly conserved in all mosquito-borne flaviviruses (Fig. 4A). In order to determine whether dengue virus 5' and 3' CS participate in the RNA-RNA complex observed in gel shift assays, we first monitored binding of the 3' SL probe to the unlabeled 5' UTR-C62 RNA carrying a deletion of 5' CS. Incubation of the probe with the 5' UTR RNA showed no complex formation (Fig. 4B), suggesting that base pairing between 5' CS and 3' CS was necessary for the interaction. To further confirm this observation, specific mutations were incorporated into the 5' UTR-C62 RNA molecule. Thus, we generated RNAs with 2-, 4-, 5-, and 7-nucleotide substitutions within the 11-nucleotide sequence of dengue virus 5' CS (Fig. 4C, mutants 143, 144, 145, and 146, respectively). The WT 3' SL probe was incubated with different concentrations of unlabeled WT and mutated 5' UTR-C62 RNAs. The RNA binding assays indicated that a 2-nucleotide mismatch between 5' and 3' CS complementarity (mutant 143) was sufficient to decrease
the RNA binding affinity more than 10-fold, changing the $K_d$ from 8 to 100 nM (Fig. 4D). Furthermore, point mutations that included more than four base mismatches between the 5′ and 3′ CS drastically decreased RNA-RNA complex formation (Fig. 4D, mutants 144, 145, and 146). These results indicate that sequences within 5′ CS are necessary for RNA-RNA complex formation.

Mapping RNA-RNA contacts between 5′- and 3′-end sequences of dengue virus. To evaluate all possible contacts between the 5′ and 3′ ends of dengue virus RNA, we examined the predicted secondary structures of a molecule containing both the 160 nucleotides of the 5′ end (5′ UTR-C62) and the 106 nucleotides of the 3′ end of dengue virus genome by using the Mfold program (65). Two alternative folding predictions...
were obtained with similar energies, and both structures showed base pairing between 5′ and 3′ CS (Fig. 5). Similar predictions were previously reported for dengue virus and other flaviviruses (38, 52, 62). In structure I, the highly conserved 3′ SL structure was retained, and the interactions between 5′ and 3′ sequences were maintained mainly by 5′-3′ CS base pairing. Structure II is very different in that the base of the 3′ SL is disrupted, allowing base pairing to occur just upstream of the initiator AUG of the 5′ UTR (Fig. 5). This second complementary sequence of 16 nucleotides bears one mismatch and one bulge. Due to its location just 5′ to the initiator AUG, we named this second putative cyclization element 5′ UAR (upstream AUG region). To evaluate the importance of 5′-3′ UAR base pairing in RNA-RNA complex formation, we performed gel shift assays using the 5′ UTR-C62 RNAs carrying specific mutations in UAR. Because the 5′ UAR is located within a very conserved stem-loop structure (Fig. 6A), the mutations were incorporated into the loop and both sides of the stem to avoid altering the predicted stem-loop structure. Thus, we generated 5′ UTR-C62 mutants 131, 133, 175, and 170, carrying one to four substitutions, respectively (Fig. 6A). These nucleotide changes resulted in one to four mismatches within the UAR when the 5′ UTR-C62 RNA hybridizes with the 3′ SL RNA (Fig. 6B). In order to estimate the affinity between the 5′ and 3′ RNAs, the wild-type 3′ SL probe was titrated with different concentrations of each of the mutant 5′ UTR-C62 RNAs. A single G/C substitution in the 5′ UAR (mutant 131) greatly decreased the affinity for the 3′ SL RNA (Fig. 6C). Moreover, substitutions in the 5′ UAR with three or four mismatches almost abolished RNA-RNA interaction. The lack of binding of the 3′ SL RNA to RNA molecules carrying substitutions within the 5′ UAR indicates that 5′-3′ UAR interaction is involved in complex formation, suggesting that structure II (depicted in Fig. 5) is the one predicted in the RNA-RNA complex. Furthermore, these results were confirmed by AFM data, in which analysis of RNA molecules of 2.3 kb with deletions at the 3′ end including the 3′ UAR did not acquire circular conformation (data not shown).

**Significance of long-range RNA-RNA interactions for viral viability.** Several reports have previously demonstrated the requirement of 5′ and 3′ CS for viral replication in different flaviviruses (11, 38, 39, 42). Here, we demonstrated that in addition to 5′-3′ CS complementarity, a second RNA region present at the 5′ and 3′ UTR of dengue virus (named 5′ UAR and 3′ UAR) is required for RNA-RNA complex formation (see structure II in Fig. 5). Thus, to further substantiate the notion that genome circularization is indeed required for viral replication, we asked whether 5′-3′ UAR complementarity is necessary during the viral life cycle. To this end, we constructed recombinant dengue virus RNA carrying substitutions in the 5′ or 3′ UAR and in both 5′-3′ UAR which will disrupt or reconstitute the 5′-3′ complementarity, respectively. The 5′ UAR mutant contained three nucleotide substitutions (80-AC AAGACAAGUG [the substitutions are underlined]) (Fig. 7A, DV-5′ UAR 175). This mutation was shown to abolish RNA-RNA complex formation in the RNA binding assay (Fig. 6, mutant 175). The recombinant viral RNA carrying the mutation at the 3′ end contained four substitutions (three substitutions in the 3′ UAR [underlined] [10634-CAACAGAUCGU GCUGUGU] and one G/C substitution at position 10718).

The three substitutions in the UAR were designed to restore complementarity with mutant 5′ UAR 175, and the fourth substitution was designed to maintain the predicted structure of the 3′ SL (Fig. 7A, DV-3′ UAR 177). We also generated the recombinant dengue virus RNA containing both mutations at the 5′ and 3′ ends in a single RNA (DV-5′-3′ UAR 175-177) simultaneously.

In order to confirm that the mutations designed at the 5′ and 3′ UAR based on RNA-rolling predictions indeed restore RNA-RNA interaction, we performed gel shift analysis with a 3′ SL probe carrying the mutation in the 3′ UAR (3′ SL 177). This probe was incubated with unlabeled WT or mutated 5′ UTR-C62 RNA carrying mutation 175. The RNA binding assay indicated that the 3′ SL 177 probe interacted very weakly with the WT 5′ UTR-C62 RNA, but RNA-RNA binding affinity was greatly increased when mutant 175 was used as the unlabeled RNA (Fig. 7B, lanes 10 to 17). Therefore, we concluded that combining the mutations in the 5′ and 3′ UAR restores RNA-RNA interaction. Titrations of the probes 3′ SL WT and 3′ SL 177 with different concentrations of unlabeled RNAs indicated that the affinity between the two mutant RNAs is similar to that observed with the two WT RNAs (Fig. 7B, compare lanes 1 to 4 with lanes 10 to 13).

Next, we in vitro transcribed and transfected into BHK cells full-length dengue virus RNAs corresponding to the WT and mutants DV-5′ UAR 175, DV-3′ UAR 177, and DV-5′-3′ UAR 175-177. Initially, the infectivity of the RNAs was assessed by IFA for dengue virus antigens in transfected cells by using murine anti-dengue virus type 2 antibodies. The cells transfected with DV-5′ UAR 175 and DV-3′ UAR 177 RNAs were negative at 3, 6, 9, 12, 15, 18, and 21 days (Fig. 7C). For this experiment, cells were reseeded at 3-day intervals. In contrast, when the cells were transfected with the double mutant DV-5′-3′ UAR 175-177 RNA, the IFA was positive at day 3 after transfection, and about 10% of the monolayer was positive at day 9 (Fig. 7C). The WT RNA-transfected cells were positive for dengue virus antigens by IFA by 24 h, and nearly 100% of cells in the monolayer were positive at day 4. This result indicates that mutations in both the 5′ and 3′ UAR that reconstitute RNA-RNA interactions rescue the lethal phenotype of substitutions either at the 5′ or the 3′ UAR sequences. Assuming that the efficiency of transfection of cells with WT and mutant RNAs was similar, this result indicates that the DV 5′-3′ UAR 175-177 mutant virus did replicate in the transfected cells but that replication was markedly impaired in comparison with that of the WT virus.

To confirm these results, we searched for infectious dengue virus particles after transfections. The media from cells transfected with WT and mutated RNAs were collected after 3, 6, 9, 12, 15, 18, and 21 days and used for plaque assays in BHK cells. Infectious particles were not recovered for the single mutants at the 5′ and 3′ UAR. In contrast, the double-mutant RNA yielded viral particles with a small-plaque phenotype in comparison to that of WT virus (Fig. 7D). The plaque size observed after 9 days of infection of BHK cells with the mutant virus collected after 9, 12, 15, 18, and 21 days after transfection remained small (2 mm compared with 4 mm of the WT). Furthermore, we analyzed whether nucleotide changes in the mutant viruses occurred during virus cultivation. To this end, RNA was extracted from viruses collected in the media from...
A

Dengue Virus Genome

WT  DV 5'UAR Mut 175  WT  DV 3'UAR Mut 177

B

5'UTR-C62 RNA (nM)

<table>
<thead>
<tr>
<th></th>
<th>WT</th>
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<td>20</td>
<td>5</td>
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RNA-RNA complex  3'SL Probe

1  2  3  4  5  6  7  8  9  10  11  12  13  14  15  16  17

Probe 3'SL WT  Probe 3'SL 177

C

Day 3  Day 6  Day 9

DV 2 WT  DV 5'UAR Mut 175  DV 3'UAR Mut 177

DV 5'3'UAR Mut 175-177

D

DV 2 WT  DV 5'3'UAR Mut 175-177
transfected cells at different times and used for RT-PCR. Sequencing analysis of the cDNAs revealed that the original substitutions in the double-mutant virus were retained. These results were reproduced in three independent experiments.

Taken together, these results indicate that point mutations in either the 5' or the 3' UAR were lethal and that reconstitution of the RNA-RNA complementarity between these sequences was essential for viral viability. In addition, the impaired replication of the virus DV-5'3'UAR 175-177 strongly suggested that UAR nucleotide sequence and/or RNA structures involving the UAR at the 5' UTR and the 3' SL is required for efficient dengue virus replication.

**DISCUSSION**

In this study, we provide direct evidence for long-range RNA-RNA interactions in the dengue virus genome. Using AFM, we visualized individual RNA molecules in circular conformations. Novel complementary sequences present at the 5' and 3' ends of the viral genome were identified as determinants for RNA-RNA contacts. More importantly, analysis of recombinant dengue viruses indicate that complementarity between these sequences is essential for viral replication. The data presented here, together with previous reports, support the notion that communication between 5' and 3' ends of RNA molecules could be a common feature of both host mRNAs and viral RNA genomes (6, 30–32, 59).

AFM is a powerful technique that has been shown to be useful for studying conformations of DNA, structures of DNA-protein complexes, and motion of enzymatic reactions (such as transcription) at the single-molecule level (9, 24, 26, 28, 29, 36, 45). In contrast, fewer studies analyzing biologically relevant RNA molecules have been reported (1, 2, 53, 59). Here, we used AFM to investigate long-range RNA-RNA interactions between conserved RNA elements present in the genome of mosquito-borne flaviviruses. RNA molecules of 2.3 kb carrying dengue virus sequences at the 5' and 3' ends acquire circular conformations (Fig. 2), while control RNA with deletions of specific dengue virus sequences at the 3' end was unable to form circles (Table 1). Similar results were obtained using the full-length dengue virus RNA of 10.7 kb (Fig. 3). The statistically significant amount of circular molecules found in the samples indicated that the affinity between the RNA elements present at the ends of the dengue virus genome was sufficient for RNA circularization. Analysis of RNA-RNA complexes formed in vitro together with AFM data indicated that at least two pairs of complementary sequences were necessary for RNA-RNA interactions. We demonstrated that base pairing between the 5' and 3' CS alone was not sufficient for in vitro RNA-RNA complex formation. A second complementary sequence present at the 5' end just upstream of the initiator AUG and at the 3' end within the 3' SL (5' and 3' UAR) was also required. Single nucleotide changes within the CS or UAR showed a large effect on the affinity of the 5' and 3' RNAs (Fig. 4 and 6), suggesting that interactions between the two pairs of complementary sequences are important to stabilize the RNA-RNA association. However, the absolute requirement of Mg2+ for complex formation strongly suggests that 5'–3' association also involves tertiary structures in the RNA.

The 5' and 3' CS are 10 or more contiguous nucleotides that complement perfectly in all mosquito-borne flaviviruses. Sequence analysis indicates that at least eight of these nucleotides are identical in these viruses (Fig. 4A), suggesting that even though coevolution of the two complementary sequences could occur, there must be a growth advantage to preserve the nucleotide sequences. A number of reports using mutational analysis in different flaviviruses have demonstrated the requirement of 5' and 3' CS for viral replication (11, 38, 39, 42). In contrast, a possible role of 5' and 3' UAR sequences in viral replication was not previously inspected. Here, we show that specific substitutions within the 5' or 3' UAR in the context of infectious dengue virus type 2 yielded no viable viruses. Importantly, mutations at the 5' and 3' UAR that restore complementarity were sufficient to rescue viral replication, demonstrating that base pairing of 5'-3' UAR provides an essential element for viral viability. In addition, the small-plaque phenotype observed with the mutant virus suggests that the nucleotide sequence of the 5' and 3' UAR is required for efficient viral replication. Because the cyclization motives are widely conserved among flaviviruses, the results presented here could be extrapolated to other members of this genus.

**Roles of RNA cyclization during viral replication.** It is a common notion that RNA elements at the 5' end of viral RNAs modulate the efficiency of translation initiation, while elements at the 3' end recruit the replication machinery to initiate negative-strand RNA synthesis. This view is now...
changing, as accumulating evidence indicates that the roles of both the 5' and 3' ends functionally overlap. For instance, sequences at the 5' end of poliovirus, Sindbis virus, and alfalfa mosaic virus are essential for negative-strand synthesis at the 3' end (16–19, 21, 55), while cis-acting elements at the 3' UTR of viral genomes have been shown to modulate the efficiency of translation initiation at the 5' end (3, 44, 57, 58). In dengue virus, it was proposed that sequences within the 3' CS annealed to the 3' CS to provide the recognition signals for RNA synthesis by the viral NS5 polymerase (62, 63). Cyclization of the dengue virus genome is likely to induce changes both upstream of the initiator AUG and within the structure of the very conserved 3' SL (Fig. 5). Because these two elements are cis-acting signals for translation and RNA synthesis, it is possible that changes in the RNA conformation could modulate these steps of viral replication. Taken together, we hypothesize that bringing the initiation sites for translation and RNA synthesis physically together provides a strategy to coordinate both processes. Further studies using dengue virus replicas expressing a reporter gene are under way in our laboratory to dissect the role(s) of circular conformations of the RNA in each step of the viral life cycle.

In the present study, we observed RNA-RNA interactions in the absence of proteins. However, it is likely that cellular or viral proteins modulate the formation or stabilization of circular or linear RNAs in vivo. In fact, it has been proposed that the viral protease NS3 binds to dengue virus 3' SL (12), the cellular protein EF-1α interacts with several flavivirus 3' SLs (5, 13), and the cellular La protein binds both 5' and 3' UTR elements (20). In the case of Japanese encephalitis virus, both NS3 and NS5 proteins bind cooperatively to the 3' SL (10). Because the cyclization 3' UAR sequences are located within the 3' SL structure, binding of cellular or viral proteins to this RNA element could disrupt or enhance cyclization of the viral genome in the infected cell.

Clearly, secondary and tertiary structures of viral RNAs serve highly conserved functions in viral replication. In addition, these structures of the RNA could change during the viral life cycle, adopting different conformations during translation, RNA synthesis, and encapsidation. Therefore, further analysis of RNA conformations acquired during these processes will help to clarify molecular details of viral replication.

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CIRCULARIZATION OF FLAVIVIRUS RNA


