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Reprint of: Core protein-mediated 5′–3′ annealing of the West Nile virus genomic RNA in vitro☆

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Genome cyclization through conserved RNA sequences located in the 5′ and 3′ terminal regions of flavivirus genomic RNA is essential for virus replication. Although the role of various cis-acting RNA elements in panhandle formation is well characterized, almost nothing is known about the potential contribution of protein cofactors to viral RNA cyclization. Proteins with nucleic acid chaperone activities are encoded by many viruses (e.g., retroviruses, coronaviruses) to facilitate RNA structural rearrangements and RNA–RNA interactions during the viral replicative cycle. Since the core protein of flaviviruses is also endowed with potent RNA chaperone activities, we decided to examine the effect of West Nile virus (WNV) core on 5′–3′ genomic RNA annealing in vitro. Core protein binding resulted in a dramatic, dose-dependent increase in 5′–3′ complex formation. Mutations introduced in either the UAR (upstream AUG region) or CS (conserved sequence) elements of the viral RNA diminished core protein-dependent annealing, while compensatory mutations restored the 5′–3′ RNA interaction. The activity responsible for stimulating RNA annealing was mapped to the C-terminal RNA-binding region of WNV core protein. These results indicate that core protein – besides its function in viral particle formation – might be involved in the regulation of flavivirus genomic RNA cyclization, and thus virus replication.

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1 Introduction

Flaviviruses constitute a large and diverse group within the Flaviviridae family of positive-strand enveloped RNA viruses (Lindenbach et al., 2007). Medically important flaviviruses include the mosquito-borne West Nile virus (WNV), yellow fever virus (YFV), Japanese encephalitis virus (JEV), and dengue virus (DENs serotypes 1–4), as well as tick-borne encephalitis viruses (TBEVs). Infection by arthropod-borne flaviviruses is associated with significant morbidity and mortality worldwide (Gould and Solomon, 2008; Griffin, 2011; Guzman et al., 2010). Increased urbanization, intercontinental travel, failure of vector mosquito control and increasing global temperatures have collectively resulted in the recent emergence or re-emergence of mosquito-borne flaviviruses in previously disease-free areas (Gould and Higgs, 2009; Kilpatrick, 2011). West Nile virus first appeared in the New World in 1999, causing a viral encephalitis outbreak in New York City (Petersen, 2009). Since then, the virus has rapidly spread throughout the US, Central and South America, reaching Argentina by 2006 (Petersen, 2009). Due to the increased geographic dispersal of the Asian tiger mosquito Aedes albopictus, dengue virus is currently re-emerging in Southern Europe (Gjenero-Morgan et al., 2010; La Ruche et al., 2010), after almost a hundred-year hiatus (Rosen, 1986).

Flaviviruses are small, enveloped viruses with a single-stranded, positive sense RNA genome of ~11 kb. A single open reading frame (ORF) encodes a large polyprotein precursor which is co- and post-translationally processed to yield the viral structural proteins (C-prM-E), and the non-structural proteins (NS1–NS2A–NS2B–NS3–NS4A–NS4B–NS5) directing genome replication (Lindenbach et al., 2007). The ORF is flanked by a short (~100 nt) 5′ untranslated region (UTR) and a longer (~400–800 nt) 3′ UTR, both of which contain highly conserved RNA secondary structures and RNA sequences involved in the regulation of viral translation and RNA replication [reviewed in (Brinton, 2002; Markoff, 2003); see Fig. 1].
The 5' UTR begins with a Y-shaped stem-loop (SLA) (Brinton and Dispoto, 1988) necessary for viral replication (Filomatori et al., 2006). The initiator AUG codon for polypeptide synthesis is situated in a second stem-loop structure (SLB), which is followed by the capsid-coding region hairpin (CHP) regulating start codon selection (Clyde et al., 2008; Clyde and Harris, 2006). The 3' UTR consists of a variable region, followed by tandem dumbbell-like structures (DB1 and DB2), and a highly stable and conserved terminal stem-loop (3' SL) (Brinton et al., 1986). In addition, a complex network of pseudoknot interactions is thought to influence the topology and function of the 3' UTR (Funk et al., 2010; Olsthoorn and Bol, 2001; Pijlman et al., 2008; Shi et al., 1996; Silva et al., 2010).

The positive-sense genomic RNA of flaviviruses is first copied by the viral RNA-dependent RNA polymerase (RdRp) to generate a complementary minus-strand RNA that, in turn, serves as a template for the amplification of plus-strand viral genomes. Interestingly, in vitro polymerase assays demonstrated that minus-strand RNA synthesis requires the presence of both the 5' and 3' UTRs (You et al., 2001; You and Padmanabhan, 1999). The promoter element for minus-strand RNA synthesis was later identified as stem-loop A (SLA, Fig. 1) in the 5' UTR, which binds the viral RdRp.
with high affinity (Filomatori et al., 2011, 2006; Lodeiro et al., 2009). According to current models, based mostly on work in dengue viruses, panhandle formation through complementary sequences in the 5′-core and 3′ UTR regions would reposition the 3′ end of the genome in the vicinity of the promoter–RdRp complex, coupling transcription initiation to genome cyclization (reviewed in Gebhard et al., 2011). At the same time, the long-distance RNA interaction is believed to be accompanied by structural rearrangements in the highly conserved 3′ stem-loop (3′ SL, Fig. 1) (Dong et al., 2008), allowing RdRp binding to the terminal nucleotides of the genome (Gebhard et al., 2011). Genomic RNA cyclization in mosquito-borne flaviviruses is mediated by three pairs of long distance RNA interactions: 5′–3′ UAR (upstream AUG region) (Alvarez et al., 2005b), 5′–3′ DAR (downstream AUG region) (Dong et al., 2008; Friebe and Harris, 2010), and 5′–3′ CS (conserved sequence) (Hahn et al., 1987) (Fig. 1). Mutations disrupting complementarity in any of these elements abrogate viral replication in subgenomic replicon model systems (Alvarez et al., 2008, 2005b; Breidenbeck et al., 2003; Conver et al., 2003; Friebe and Harris, 2010; Friebe et al., 2011; Khromykh et al., 2001; Lo et al., 2003; Men et al., 1996; Villordo et al., 2010; Zhang et al., 2008), while compensatory mutations rescue viability (Alvarez et al., 2008, 2005b; Friebe and Harris, 2010; Friebe et al., 2011; Khromykh et al., 2001; Lo et al., 2003; Zhang et al., 2008).

Flavivirus genomic RNA is encapsidated by the small, highly basic core protein. Core corresponds to the N-terminus of the polyprotein, from where it is released in its mature form by the action of the NS2B-NS3 serine protease complex (Amberg et al., 1994; Lobigs, 1993; Yamshchikov and Compans, 1994). Despite relatively little sequence conservation, all flavivirus core proteins share a common functional and structural domain organization. The N- and C-terminal extremities contain a high concentration of basic amino acids, and bind to the viral genomic RNA independently (Khromykh and Westaway, 1996). Flanked by the RNA-binding regions, an internal hydrophobic domain is responsible for the dimerization/oligomerization (Bhuvanakhantham and Ng, 2005; Wang et al., 2004), as well as for the membrane/lipid association of core protein (Markoff et al., 1997; Samsa et al., 2009).

Structural studies by circular dichroism, X-ray crystallography and nuclear magnetic resonance (NMR) on DENV, WNv, and YFV core proteins (Dokland et al., 2004; Ivanji-Nagy et al., 2008; Jones et al., 2003; Ma et al., 2004) also support a conserved structure, with a highly flexible N-terminal RNA-binding region followed by three or four alpha-helices. Interestingly, deletion analyses demonstrated a remarkable functional flexibility of core protein, suggesting that precisely folded three-dimensional structures are not required for RNA binding and membrane association, and these functions are rather determined by the overall physicochemical nature of the domains. Large deletions are well tolerated in the internal hydrophobic region of both mosquito-borne and tick-borne flaviviruses (Koller et al., 2002, 2003; Schlick et al., 2009; Zhu et al., 2007). Similarly, the N- and C-terminal extremities can function (at least partly) redundantly in RNA packaging (Patkar et al., 2007).

Core proteins of WNv (Ivanji-Nagy et al., 2008), dengue virus (Pong et al., 2011), as well as cores in the related hepativiruses (Cristofari et al., 2004; Ivanji-Nagy et al., 2006) and pestiviruses (Ivanji-Nagy et al., 2008), were shown to facilitate nucleic acid rearrangements without ATP consumption, acting as efficient RNA chaperones in vitro. The RNA chaperone activity of WNv core protein was mapped to the C-terminal RNA-binding region of the protein (Ivanji-Nagy et al., 2008). In this study, we examined the effect of WNv core protein chaperoning on viral 5′–3′ UTR annealing, using an in vitro model system with separate 5′ and 3′ RNAs. We found that core protein binding greatly increases the rate of 5′–3′ complex formation, and is required for the interaction when full-length 3′ UTR RNAs are used (Fig. 2). Mutations abolishing either the UAR or CS interaction diminished, but did not completely abrogate core-protein induced annealing, while compensatory mutations restored the interaction (Fig. 3). In agreement with the results of in vitro chaperone assays (Ivanji-Nagy et al., 2008), stimulation of RNA annealing was mapped to the C-terminal RNA-binding region of core protein (Fig. 4).

2. Materials and methods

2.1. Plasmid construction

For cloning the 3′ UTR of West Nile virus (Eg101 strain, GenBank accession AF260968), total RNA was extracted from virus infected Vero cells and reverse transcribed using the ThermoScript RT-PCR system (Invitrogen). Reverse transcription was carried out at 60 °C, using ODN Eg101-3′UTR-ss as a primer (Table 1). cDNA was amplified by Eg101-3′UTR-ss and Eg101-3′UTR-as and cloned between the Sall and HindIII sites of pSP64 (Promega), resulting in pSP64-3′UTR vector. Deletions in the 3′ UTR were introduced by amplifying the desired regions in pSP64-3′UTR by PCR, and re-cloning the fragments between the Sall and HindIII sites of pSP64.

The 5′ UTR-core region (nt 1–164 of the viral genome) was cloned by annealing of overlapping ODNs (Eg101-5′UTR-D0 to -D5), followed by ligation between the Xmal and EcoRI sites of pSP64 vector (Promega).

Mutations were introduced in the UAR or CS region of pSP64-3′UTR or pSP64-5′UTR by core using a PCR-based mutagenesis protocol (Mikaelian and Sergeant, 1992), with three common ODNs (pSP64, pSP64-T7, and SP6) and one mutation-specific ODN (mut-UAR or mut-CS) for each mutant.

All plasmid constructs were verified by sequencing.

2.2. Proteins and peptides

Full length WNv core protein (amino acids 2–105; GenBank accession number AF481864) was expressed in Escherichia coli and purified as previously described (Ivanji-Nagy et al., 2008). Core peptides WNv C(1–24) and WNv C(80–105), corresponding to the N- and C-terminal RNA-binding regions of the core protein, were synthesized by Fmoc-OH/DCC/Hobt chemistry and purified as previously described (Ivanji-Nagy et al., 2008).

2.3. In vitro RNA synthesis

Plasmids containing the 5′ UTR-core region or fragments of the 3′ untranslated region of WNv strain Eg101 were linearized by digestion with Xmal or HindIII restriction enzymes, respectively. In vitro transcription was carried out using T7 RNA polymerase, according to the manufacturer’s instructions (Promega). 5′ UTR-core RNAs were labelled by incorporation of α32-P UMP during in vitro transcription. RNAs were purified on 6% denaturing polyacrylamide gels containing 7 M urea in 50 mM Tris-borate (pH 8.3)–1 mM EDTA (0.5 × TBE), and recovered by elution in 0.3 M sodium acetate–0.1% SDS overnight at 37 °C, followed by ethanol precipitation.

2.4. RNA annealing assays

In vitro synthesized RNAs were heat denatured for 2 min at 95 °C and chilled on ice. 0.1 pmol of each 5′ UTR-core and 3′ UTR RNA were mixed with annealing buffer to a final concentration of 20 mM Tris–Cl, pH 7.0, 30 mM NaCl, 0.1 mM MgCl2, 10 μM ZnCl2,
10 U RNasin and 5 mM DTT in 10 µl final volume. WNV core protein or core peptides were added to final protein to RNA nucleotide molar ratios as indicated in the figure legends (typically between 1/40 and 1/5 protein/nucleotide ratios). Reactions were incubated at 37°C for 10–15 min and quenched by adding stop solution (0.5% SDS–25 mM EDTA). Proteins were removed by proteinase K digestion and phenol–chloroform extraction. The purified RNA samples were resolved by 8% native polyacrylamide gel electrophoresis in 0.5× TBE and analysed by autoradiography and Phosphorimager quantification.

3. Results

3.1. Chaperoning 5′–3′ RNA interactions by the core protein

Cyclization of the genomic RNA is essential for viral replication in all mosquito-borne flaviviruses (Alvarez et al., 2008, 2005b; Corver et al., 2003; Khromykh et al., 2001; Lo et al., 2003; Zhang et al., 2008), but the determinants and regulation of panhandle formation are still poorly understood. Although complex formation between 5′ and 3′ UTR RNAs can be readily detected in vitro in the
absence of protein cofactors (Alvarez et al., 2008, 2005b; Villordo et al., 2010; Zhang et al., 2010), these interactions usually require high magnesium and RNA concentrations and involve short RNA molecules, thus minimizing the possibility for the RNA to become kinetically trapped in non-functional conformation(s). But in the cellular milieu, RNA folding and RNA–RNA interactions are perhaps universally facilitated by RNA chaperones and/or specific RNA-binding proteins (Cristofari and Darlix, 2002), either encoded by viruses or hijacked from the host for chaperoning viral translation, replication, and packaging.

We have previously shown that the core protein of WNV possesses potent RNA chaperone activities in vitro (Ivanyi-Nagy et al., 2008), facilitating nucleic acid annealing and RNA structural rearrangements. In order to analyse the possible effect of core protein on panhandle formation in the WNV genome, RNA molecules corresponding to nucleotides 1–164 of the genomic RNA (RNA 1–164; Fig. 2A) were in vitro synthesized, radioactively labelled and incubated with equal amounts of non-labelled 3′ FL-UTR RNA (Fig. 2A) in the presence of varying amounts of full-length WNV core protein. Following proteinase K digestion of proteins and phenol–chloroform purification of RNAs, 5′–3′ RNA complex formation was assessed by electrophoretic mobility shift assays (Fig. 2B, lanes 1–5). WNV core induced a dose-dependent increase in RNA–RNA interactions, resulting in almost complete annealing at 1 protein to 5 nt molar ratio (Fig. 2B, lane 5; compared to lane 1 in the absence of core protein). In order to examine the potential effect of intramolecular interactions on annealing in the 3′ UTR, 5′ deleted 3′ UTR RNAs, lacking either the variable region (3′ΔVR RNA), or most of the UTR except for the cyclization sequences and the 3′ stem-loop (3′CYC RNA) were examined (Fig. 2B, lanes 6–10 and 11–15, respectively). For all RNA molecules, incubation with core protein induced efficient annealing in a dose-dependent manner (Fig. 2B, lanes 7–10 and 12–15). Interestingly, core protein-independent annealing was more pronounced for the shortest 3′CYC RNA (lane 11 vs lanes 6 and 1), suggesting that RNA sequences 5′ to the cyclization signals might change the topology of the 3′ UTR and interfere with the annealing reaction. To further characterize these differences, a time-course analysis of annealing, either without core protein (Fig. 2C), or in the presence of full-length core (Fig. 2D) was carried out. Without core protein, annealing for the 3′FL-UTR RNA and 3′ΔVR RNA was hardly detectable during the 1 h incubation period, while for the shortest 3′CYC RNA, around one third of the molecules formed 5′–3′ RNA interaction (Fig. 2C and E). In contrast, core protein dramatically increased complex formation, resulting in high levels of annealing as early as 30 s (Fig. 2D and E). Indeed, based on second-order kinetics, the initial annealing rate was estimated to increase by ~500-fold in the presence of core (data not shown). Nevertheless, annealing of the longer RNAs was still significantly delayed compared to 3′CYC RNA,
suggesting that a kinetic barrier must be overcome for the reaction to proceed.

Phylogenetic analyses suggest that the current discontinuous cyclization sequence of mosquito-borne flaviviruses originated – possibly by template switching – as a perfectly complementary continuous region, conserved to this day in the tick-borne members of flavivirus genus (Gritsun and Gould, 2007). Thus, the UAR-DAR-CS region probably acts as a single regulatory sequence, where replication is determined by the overall stability of the long distance interaction between the 5′ and 3′ regions. In support of this, replication of WNV with a complete deletion of the 3′ CS can be rescued by second-site mutations stabilizing the UAR and DAR interactions (Zhang et al., 2010). In order to examine whether the core protein-dependent annealing of the 5′ and 3′ UTR RNAs depends on the UAR and/or CS interaction, mutations were introduced separately in these elements of RNA 1–164 (Fig. 3A), and cyclization with wild-type 3′FL-UTR RNA was assessed by electrophoretic mobility shift assays. Mutations in either the 5′ UAR or 5′ CS led to a significant decrease in annealing (Fig. 3B, lanes 6–10 and 11–15), while cyclization was rescued by compensatory mutations introduced in the 3′ UAR or 3′ CS of the 3′FL-UTR RNA (lanes 16–20 and 21–25, respectively). These results suggest that the core protein-dependent in vitro annealing recapitulates the features of the cyclization sequence analysed in subgenomic or genome-length cellular models. Interestingly in vitro RNA annealing was readily detectable even with 5 mutations at high core protein levels, while even single point mutations result in a lethal phenotype in subgenomic replicons. This suggests that chaperoning by core protein may stabilize the weak interaction present in the mutants, thus partially relaxing the complementarity requirements of the long distance interaction (Basu and Brinton, 2011; Zhang et al., 2010; see Section 4).

3.2. The C-terminal domain of WNV core protein chaperones 5′–3′ RNA interactions

Flavivirus core proteins have two independent RNA-binding regions at the N- and C-terminal extremities of the protein (Khromykh and Westaway, 1996; Fig. 4A). We have previously shown that the two RNA-binding domains do not act in synergy and that only the C-terminal basic region was active in in vitro nucleic acid chaperone assays (Ivanyi-Nagy et al., 2008). In order to delineate the requirements of core protein for facilitating 5′–3′ UTR interaction, synthetic peptides corresponding to the N- and C-terminal RNA-binding regions (WNV C(1–24) and WNV C(80–105), respectively; Fig. 4A) were used in the annealing reactions with wild-type RNAs. While WNV C(1–24) did not influence 5′–3′ annealing (Fig. 4B, lanes 7–12), WNV C(80–105) induced an increase in 5′–3′ complex formation (Fig. 4B, lanes 13–18). Nevertheless, at equal molar ratios, incubation with full-length core protein resulted in higher levels of complex formation than with WNV C(80–105), indicating that sequences or structural features outside the C-terminal region contribute to full chaperoning activity.

4. Discussion

Complementarity between the genome-terminal cyclization sequences, presumably resulting in panhandle formation in vivo, is absolutely required for flavivirus RNA replication (Khromykh et al., 2001). Although formation of the long-distance interaction is believed to be thermodynamically favoured, the annealing reaction is prohibitively slow (or may not even reach completion) under physiologically relevant conditions (Fig. 2). RNA chaperones are able to disrupt transient or non-functional RNA interactions,
thereby decreasing the kinetic barrier hindering the formation of the most stable RNA conformation (reviewed in Cristofari and Darlix, 2002; Schroeder et al., 2004). Indeed, RNA chaperoning might increase the speed of intramolecular RNA rearrangements or intermolecular RNA–RNA interactions several thousand-fold without ATP consumption. We have previously shown that core proteins in the Flaviviridae family, including that of WNV, hepatitis C virus and bovine viral diarrhea virus, possess potent RNA chaperone activities (Cristofari et al., 2004; Ivanyi-Nagy et al., 2006, 2008), facilitating nucleic acid annealing reactions with various DNA and RNA substrates. The molecular mechanism of chaperone action is still poorly understood, and it probably involves a combination of charge neutralization, molecular crowding, and local melting of nucleic acid structures by entropy transfer (Tompa and Csermely, 2004). High affinity binding of a protein to the substrate DNA/RNA is not sufficient, in itself, to trigger the conformational changes observed upon core protein chaperoning (Cristofari et al., 2004; Ivanyi-Nagy et al., 2008).

In this study, we analysed the RNA chaperone activity of WNV core using its cognate target molecules, corresponding to the highly structured terminal regions of the viral genomic RNA. WNV core protein was found to induce a dramatic acceleration in the 5′–3′ UTR annealing reaction (Figs. 2–4), suggesting a possible link between genome cyclization, replication, and packaging.

Paradoxically, with the exception of a short N-terminal sequence which contains the 5′ CS RNA element, the structural protein-coding region is not essential for WNV genomic RNA replication (Kromykh and Westaway, 1997). The discrepancy between the in vitro activity of core protein and its in vivo dispensability suggests that flaviviruses might hijack cellular RNA-binding proteins that may compensate for the loss of core chaperoning. Indeed, an RNA–protein–RNA co-immunoprecipitation assay identified seven distinct proteins (with molecular weights of 35, 37, 40, 45, 52, 76, and 97 kDa) interacting with both the 5′ UTR and 3′ UTR RNAs of dengue virus (Garcia-Montalvo et al., 2004). Although most of these cellular proteins remain to be identified and their interaction with the genomic RNA verified, the 52 kDa band was confirmed as being La protein, an abundant cellular RNA chaperone (Belisova et al., 2005; Chakshusmathi et al., 2003). La protein was shown to interact with the UTRs of dengue virus and Japanese encephalitis virus (De Nova-Ocampo et al., 2002; Garcia-Montalvo et al., 2004; Vashist et al., 2009, 2011; Yocupicio-Monroy et al., 2007), and to bind to the viral replication proteins NS5 and NS3 (Garcia-Montalvo et al., 2004). At present, the effect of La binding on genomic RNA cyclization is still controversial. Recombinant La protein led to decreased RNA synthesis in an in vitro viral replicase assay, leading to the suggestion that its binding inhibited RNA cyclization (Yocupicio-Monroy et al., 2007). However, co-precipitation of the 5′ and 3′ genomic RNA regions was stimulated by the presence of increasing amounts of La protein (Vashist et al., 2011). A possible explanation for these seemingly contradictory findings might be provided by the results of Villordo and co-workers, who have shown that a balance between the linear and circular conformations of the genomic RNA, rather than cyclization per se, is important for efficient viral replication (Villordo et al., 2010). Thus, overly efficient RNA cyclization, as well as the lack of it, might both be deleterious for RNA synthesis. Besides La protein, a number of other cellular RNA chaperones, including heterogeneous nuclear ribonucleoprotein A1 (hnRNP A1), hnRNP A2, hnRNP Q, Y-box binding protein (YB–1), and polyuridylic acid tract binding protein (PTB), have been described to interact with flaviviral genomic RNA (Agis-Juarez et al., 2009; De Nova-Ocampo et al., 2002; Katoh et al., 2011; Paranjape and Harris, 2007). Although the effect of these chaperones on genome cyclization is currently unknown, they might participate in the regulation of panhandle formation in the absence of (or in addition to)
core protein, thus masking the in vivo effect of core on genome replication.

Nevertheless, recent mutagenesis studies have provided indirect evidence for the involvement of the core protein region in genome circularization and RNA replication (Basu and Brinton, 2011; Zhang et al., 2010). Genome-length viral RNAs with multiple adjacent mutations in CS or with a complete deletion of the 3′ CS element were replication competent and generated revertants or second-site mutations upon passages, restoring efficient panhandle formation (Basu and Brinton, 2011; Zhang et al., 2010). In contrast, disruption of the 5′–3′ CS interaction invariably resulted in a lethal phenotype when analysed in subgenomic replicons lacking core protein (Corver et al., 2003; Khromykh et al., 2001; Lo et al., 2003). Similarly, a deletion in the core coding region of a genome-length replicon also precluded the rescue of the CS mutant (Zhang et al., 2010), although it is still debated whether this was due to the lack of core protein expression or to the absence of RNA secondary structures in the core-coding region (Friebe et al., 2012). These results, together with our findings (Fig. 3B), suggest that cellular proteins can only partially substitute for core protein chaperoning in genome cyclization, and the requirements for panhandle formation might be more relaxed in the presence of core. This is especially important since mutations in the cyclization sequences are considered for the design of live-attenuated flavivirus vaccines, with the aim to reduce the risk of recombination with Naturally circulating viruses (Suzuki et al., 2008).

The annealing efficiency of 3′ UTR RNAs was found to be highly dependent on the length of the RNA used (Fig. 2). While 3′ CIC RNA, containing the cyclization sequences and the 3′ SL structure, could interact with the 5′ region even in the absence of core protein, the longer 3′ΔVR and 3′FL-UTR RNAs required core for 5′–3′ complex formation, suggesting that the presence of the dumbbell structures (DB1 and DB2) might interfere with annealing. Pseudoknot interactions, conserved in mosquito-borne flaviviruses, have been suggested to form between the loop of the dumbbells and a single-stranded region, including the CS element, in the linear form of the viral RNA (Olsthoorn and Bol, 2001; Figs. 1 and 2). Total or partial deletion of the individual dumbbell structures was found to seriously compromise RNA replication (Alvarez et al., 2005a; Manzano et al., 2011; Men et al., 1996), resulting in attenuated viruses that are actually pursued as a vaccine candidate for dengue viruses (Durbin et al., 2001; Troyer et al., 2001; Whitehead et al., 2003). Our results suggest that core protein chaperoning might be required to resolve the pseudoknots, yielding an RNA conformation competent for 5′–3′ annealing. Thus, the pseudoknot structures may constitute an additional layer of regulation in genome cyclization, contributing to the delicate balance between the linear and circular RNA forms.

In addition to the genomic RNA, a small subgenomic RNA (sRNA), corresponding to the last 300–500nts of the positive-strand viral RNA, also accumulates in cells infected by flaviviruses (Durbin et al., 2004; Pijman et al., 2008; Urosevic et al., 1997). sRNA is generated by the incomplete degradation of the viral genome by the host exonuclease XRN1, where the 3′ RNA region is protected by a preserved pseudoknot structure (Funk et al., 2010; Pijman et al., 2008; Silva et al., 2010). Kunjin virus sRNA was not required for viral replication but was found to be essential for cytopathicity in cell culture and for pathogenicity in infected mice (Pijman et al., 2008). As sRNA contains all the RNA elements required for interaction with 5′ cyclization sequences, its accumulation might regulate (–) strand RNA synthesis by inhibiting panhandle formation or by sequestering host or viral proteins (including core protein) interacting with the 3′ UTR (Fan et al., 2011). The interaction of core protein with sRNA and the possible consequences remain to be experimentally verified.

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