Crystal structure of a tick-borne flavivirus RNA-dependent RNA polymerase suggests a host adaptation hotspot in RNA viruses

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ABSTRACT

The RNA-dependent RNA polymerases (RdRPs) encoded by RNA viruses represent a unique class of nucleic acid polymerases. RdRPs are essential in virus life cycle due to their central role in viral genome replication/transcription processes. However, their contribution in host adaption has not been well documented. By solving the RdRP crystal structure of the tick-borne encephalitis virus (TBEV), a tick-borne flavivirus, and comparing the structural and sequence features with mosquito-borne flavivirus RdRPs, we found that a region between RdRP catalytic motifs B and C, namely region B-C, clearly bears host-related diversity. Inter-virus substitutions of region B-C sequence were designed in both TBEV and mosquito-borne flavivirus RdRPs, we found that a region between RdRP catalytic motifs B and C, namely region B-C, clearly bears host-related diversity. Inter-virus substitutions of region B-C sequence were designed in both TBEV and mosquito-borne flavivirus RdRPs, we found that a region between RdRP catalytic motifs B and C, namely region B-C, clearly bears host-related diversity. Inter-virus substitutions of region B-C sequence were designed in both TBEV and mosquito-borne Japanese encephalitis virus backbones. While region B-C substitutions only had little or moderate effect on RdRP catalytic activities, virus proliferation was not supported by these substitutions in both virus systems. Importantly, a TBEV replicon-derived viral RNA replication was significantly reduced but not abolished by the substitution, suggesting the involvement of region B-C in viral and/or host processes beyond RdRP catalysis. A systematic structural analysis of region B-C in viral RdRPs further emphasizes its high level of structure and length diversity, providing a basis to further refine its relevance in RNA virus-host interactions in a general context.

INTRODUCTION

Flaviviruses are a diverse group of positive-strand RNA viruses with a wide geographic distribution. The majority of flaviviruses are transmitted between hematophagous arthropods and vertebrate hosts, causing human encephalitis or hemorrhagic diseases in severe cases. To date, 53 flavivirus species were listed in the Flavivirus genus of the International Committee on Taxonomy of viruses (ICTV) collection (https://talk.ictvonline.org). With respect to the transmission mode, ICTV-enlisted flaviviruses can be divided into three categories: mosquito-borne, tick-borne, and vertebrate-specific (1–3). Well-known mosquito-borne flaviviruses include yellow fever (YFV), Japanese encephalitis (JEV), dengue (DENV), West Nile (WNV) and Zika (ZIKV) viruses, which are human pathogens of global concern (4–7). Tick-borne flaviviruses associated with severe human diseases include tick-borne encephalitis (TBEV), Omsk hemorrhagic fever (OHFV), and Kyasanur Forest disease (KFDV) viruses (6,8–10). Vertebrate-specific flaviviruses, also known as No Known Vector (NKV) flaviviruses, are represented by Modoc virus (MODV) isolated from rodents and Rio Bravo virus (RBV) isolated from bats (11). Flaviviruses not yet included in the ICTV collection contain a group of insect-specific viruses mostly associated with mosquitoes, such as cell fusing agent virus (CFAV) and Kamiti River virus (KRV) (12–14).

TBEV belongs to the tick-borne flaviviruses, which are sub-divided into a mammalian group and a seabird group. TBEV is classified as one of the species within the mammalian group and forms the tick-borne encephalitis (TBE) serocomplex together with KFDV, OHFV, Langat virus (LGTV), etc. The TBEV species is divided into three
sub-types, namely Western European, Siberian and Far Eastern. Similar to other flaviviruses, TBEV possesses a single-stranded, positive-sense RNA genome of ~11 kilobases (kb). The genome contains a single open reading frame (ORF) that is flanked by 5′ and 3′ untranslated regions (UTRs) (15,16). The ORF encodes a large polyprotein of about 3400 amino acids that is cleaved, through proteolytic processing, into three structural proteins (capsid-C, premembrane/membrane-prM/M, envelope-E) and seven nonstructural proteins (NS1, NS2A, NS2B, NS3, NS4A, NS4B and NS5). NS5 is the largest and most conserved protein encoded by flaviviruses, which is a unique natural fusion of an N-terminal methyltransferase (MTase) and a C-terminal RNA-dependent-RNA polymerase (RdRP) connected by a 10-residue linker (17,18). The NS5 MTase possesses guanilyltransferase (GTase) activity in the formation of the G5′-ppp-5′A linkage, and can catalyze both guanine N7 and 2′-O methylation steps in the capping process (19,20). So far, the crystal structure of the about 260-residue NS5 MTase has been reported in more than ten flaviviruses including the meaban virus from the seabird group of the tick-borne category (19,21). The NS5 RdRP is the central component of the viral genome replication and transcription machinery. However, crystal structures are only available for several mosquito-borne flaviviruses NS5 RdRPs (WNV, DENV, JEV, ZIKV and YFV) (17,20). In recent years, a group of invertebrate-associated flavivirus segments RNA viruses have been identified (22). Among these so-called Jingmenviruses, some tick-associated species have been found capable of causing human infection and diseases (22–25). While the structural proteins of these viruses do not exhibit apparent homology with the flavivirus counterparts, the nonstructural proteins NSP1 and NSP2 show high-levels of similarity with the flavivirus NS5 and NS3, respectively (26). However, the structural information of the Jingmenvirus NSP1 remains unclear.

With an aim to understand the RdRPs of the tick-associated flavivirus and related viruses, here we report the crystal structures of the TBEV NS5 MTase and RdRP. While the overall structures are consistent with known flavivirus MTase and RdRP structures in isolation, we found that the region connecting the RdRP catalytic motifs B and C was of particular interest. This inter-motif region B–C exhibits structure and length diversity in not only flaviviruses but also in RNA viruses in general. While manipulation in this region did not have essential impact on the enzymatic activities of flavivirus RdRPs and did not abolish RNA replication, it showed a lethal phenotype in cell-based virus production experiments. We therefore propose that this region may be a host adaptation ‘hotspot’ in RNA viruses, thus enriching the understandings of the functional diversity of viral RdRPs.

**MATERIALS AND METHODS**

### Cloning and protein production

The wild type (WT, full-length) TBEV NS5 gene within the DNA clone of tick-borne encephalitis virus strain WH2012 (Far Eastern subtype, GenBank accession no. KJ755186.1) was cloned into a pET26b vector. The MTase (residues 7–264) and RdRP (residues 265–891) production plasmids were made using a site-directed, ligase-independent mutagenesis method (SLIM) with WT pET26b-TBEV-NS5 as the parental plasmid (27,28). Four full-length NS5 mutant plasmids, TBEV SS (short substitution), TBEV LS (long substitution) and JEV SS/LS, with inter-species substitutions, were made using the SLIM method and corresponding full-length constructs as the parental plasmid (17,29). The JEV NS5 variants were produced as an ubiquitin fusion protein and were subsequently processed *in vivo* by a coexpressed Ubp1 protease as described previously (17,30). Plasmids were transformed into *Escherichia coli* strain BL21-CodonPlus(DE3)-RIL or BL21(DE3)-pCIG1 for TBEV or JEV NS5 overexpression, respectively (30). Cells growth and protein induction procedures were as previously described except that for TBEV NS5 and its variants the temperature for overnight culture growth was 30°C, the post-induction temperature was 16°C, and induction time was 20 h (17).

Cell lysis, protein purification, and protein storage for full-length, MTase and RdRP constructs were performed as previously described for the JEV NS5 study (17), except that the final protein samples were stored in a buffer with higher NaCl concentration (500 mM) and 10% (v/v) glycerol for TBEV RdRP crystallization (17). The molar extinction coefficients for the NS5 constructs were calculated based on protein sequence using the ExPASy ProtParam program (http://www.expasy.ch/tools/protparam.html). The yield was in the range of 3–58 mg (lowest: full-length TBEV NS5; highest: TBEV MTase) of pure protein per liter of bacterial culture.

### Protein crystallization and crystal harvesting

Lamellar or prismatic crystals of TBEV MTase were obtained within 2 weeks by sitting drop vapor diffusion at 16°C using 8–10 mg/ml protein. Normally, 0.5 μl protein was mixed with 0.5 μl reservoir solution. Best MTase crystals were obtained in 0.085 M HEPES (pH 7.5), 8.5% (w/v) PEG8000, 15% (v/v) glycerol. TBEV RdRP crystal screening was performed at 16°C and 5°C by sitting-drop vapor diffusion, and 0.5 μl of protein solution was mixed with 0.5 μl of reservoir solution. Micro-crystals appeared under several HEPES-containing conditions. The conditions were optimized, and diffraction-quality crystals were obtained in 0.1 M HEPES (pH 7.0), 0.2 M sodium thiocyanate, 15% (v/v) pentaerythritol propoxylate (5/4 PO/OH) and 5 mM MgSO₄. Best diffracting crystals were obtained by crystal dehydration. MTase and RdRP crystals were flash-cooled and stored in liquid nitrogen prior to data collection.

### Structure determination and analyses

Single crystal X-ray diffraction data were collected at the Shanghai Synchrotron Radiation Facility (SSRF) beamline BL17U1 (wavelength = 0.9792 Å, temperature = 100 K). At least 180° of data were typically collected in 0.2 oscillation steps. The reflections of the MTase and RdRP data were integrated, merged and scaled using XDS and HKL2000, respectively (Table 1) (31,32). The initial structure solutions of the MTase and RdRP were obtained using the molecular replacement program Phaser using coordinates derived...
Enzyme-based preparation of the 33-nucleotide (nt) RNA template (T33) was as previously described (37,38). The 30-nt template (T30) and 10-nt RNA primer (P10) were chemically synthesized (Integrated DNA Technologies) and the T30 was purified as previously described (39). For primer-dependent polymerase assays, the annealing procedures of the T33/P10 construct were as previously described (38). For dinucleotide-driven polymerase assays, T30 was annealed with a GG dinucleotide primer bearing a 5′-phosphate (P2, Jena BioSciences) at a 1:5 molar ratio following previously described annealing procedures (39).

### In vitro polymerase assays

For the primer-dependent polymerase assay, a 20-μl reaction mixture containing 6 μM NS5 or its variant, 4 μM T33/P10 construct, 50 mM MES (pH 6.5), 20 mM NaCl, 5 mM MnCl₂, 2 mM MgCl₂, 5 mM tris (2-carboxyethyl) phosphine (TCEP), 300 μM ATP, and 300 μM UTP, 0.01% (v/v) nonidet P 40 (NP40) was incubated at 40°C for 1, 2, 5 min before being quenched with an equal volume of stop solution (95% (v/v) formamide, 20 mM EDTA (pH 8.0), and 0.02% (w/v) xylene cyanol).

For the P2-driven polymerase assay, a 20-μl reaction mixture containing 6 μM NS5 or its variant, 4 μM T30/P2 construct, 50 mM Tris–HCl (pH 7.5), 20 mM NaCl, 5 mM MnCl₂, 2 mM MgCl₂, 5 mM TCEP, 300 μM ATP and 300 μM UTP, 0.01% (v/v) n-dodecyl-β-D-maltoside (DDM) was incubated at 30°C for 15, 30, 45 or 60 min. The reaction was quenched by an equal volume of stop solution and the mixture was heated at 100°C for 45–60 s. The procedures for denaturing polyacrylamide gel electrophoresis (PAGE), gel staining, and quantitation were as previously described (38).

### Cells and antibodies

Baby hamster kidney cells (BHK-21) (American Type Culture Collection (ATCC), CCL-10) were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 100 units/ml of penicillin and 100 μg/ml of streptomycin. C6/36 cells were grown in RPMI 1640 medium supplemented with 10% FBS, 100 units/ml of penicillin, and 100 μg/ml of streptomycin. The BHK-21 cells and C6/36 cells were maintained in 5% (v/v) CO₂ at 37°C and 28°C, respectively. By immunizing rabbits with purified TBEV NS5 protein, rabbit-derived polyclonal antibodies against TBEV NS5 protein in the antiserum were collected and purified from immunized rabbit to detect the production of TBEV protein. Monoclonal antibody (4G2) against JEV envelope protein (E) was used to detect viral protein production of JEV. Fluorescein isothiocyanate (FITC)-conjugated goat anti-mouse IgG was used as secondary antibody.
Construction of mutant genome-length cDNA and/or replicon cDNA of TBEV and JEV

Using the full-length infectious clones of the TBEV WH2012 strain and pACYC-JEV-SA14 and the related TBEV replicon as the backbones (40), the TBEV-SS, JEV-LS, JEV-SS mutant clones were constructed. The NS5 fragments bearing the substitution were generated by overlapping PCR and were introduced to the full-length TBEV and JEV clones through XbaI/XhoI and XbaI/SalI restriction sites, respectively. All constructs were verified by DNA sequencing.

Immunofluorescence assays

Plasmids carrying TBEV WT and SS genomic and related sequences were transfected into BHK-21 cells using the Lipofectamine 3000 Reagent according to the manufacturer’s protocols. Three days post transfection, the supernatant was collected and then used to infect BHK-21 cells. Cells were collected at 1, 2, 3 days post infection and were fixed with cold 4% paraformaldehyde at room temperature (r.t.). Virus replication was detected using anti-NS5 rabbit-derived polyclonal antibodies and FITC-conjugated goat anti-rabbit IgG as primary and secondary antibodies, respectively. The nuclei were stained with hoechst33258 (Beiyotime Biotechnology) staining solution.

The infectious clones of JEV (WT, LS and SS) were linearized with XhoI and then subjected to in vitro transcription using a T7 in vitro transcription kit (Thermo Fisher Scientific). Approximately 1 µg of transcribed recombinant genomic RNAs were transfected into BHK-21 cells (mammalian) and C6/36 (mosquito) with reagent DMRIE-C (Invitrogen). Then the cell slides were fixed in cold (–20°C) 5% (vol. to vol.) acetone in methanol at r.t. for 10 min. After washing three times with PBS (pH 7.4), the fixed cells were subjected to an immunofluorescence assay (IFA) with 4G2 monoclonal antibody to examine JEV envelope (E) production.

Luciferase assay

According to the manufacturer’s protocol, the TBEV replicon and related mutant plasmids were transfected into three 24-well plates using the Lipofectamin 3000 Reagent. Samples were collected at 1, 2 and 3 days time points. The transfected BHK-21 cells were lysed with 100 µl Lysis Buffer (Renilla-Glo Luciferase Assay System from Promega), and stored at −80°C for subsequent luciferase activity analysis. After collecting all samples, the Renilla luciferase (RLuc) activity was measured by mixing 10 µl of cell lysate with 50 µl of substrate solution. All luciferase assays were performed in triplicates.

RT-qPCR analysis

According to the manufacturer’s protocol, RNA was extracted from BHK-21 cells and their supernatants using the TRIzol Reagent (Invitrogen), and then the Tiangen’s FastKing gDNA Dispelling RT SuperMix reverse transcription kit was used to reverse transcribe 1 µg of total RNA into cDNA. The cDNA was amplified following a SYBR Green-based quantitative PCR protocol (Bio-Rad) using a real-time quantitative PCR system (Bio-Rad CFX) that requires an activation at 95°C for 3 min, followed by 40 amplification cycles, with 10 s at 95°C followed by 45 s at 60°C for each cycle. Amplification in a universal pair of primers (TBEV qPCR-F: gagaagaagaaacaggtgta and TBEV qPCR-R: cccatacctctgtgtc) were used to amplify the region spanning nucleotides 2898–3013 of TBEV (corresponding to a segment in the NS1 gene), which is conserved in all TBEV strains. The RNA copies were calculated based on the standard curve composed of 10-fold serial dilutions of the standard infectious clone plasmid DNA.

Statistical analysis

Data of the luciferase assay and RT-qPCR are presented as means ± standard deviations (SD). Student’s t-test was used to assess statistical significance, with significance defined by P value <0.05 (*), <0.01 (**) or <0.001 (***)

RESULTS

The overall structure of TBEV NS5 MTase and RdRP are largely consistent with known flavivirus NS5 structures

With an aim to understand the structural features of NS5 RdRP from tick-borne flaviviruses and related species, we attempted to crystallize the full-length TBEV NS5 but did not obtain any hits in the screening. Truncated forms of TBEV NS5 were also tested in crystallization trials. Two of these forms, containing the MTase and RdRP modules, respectively, were crystallized, yielding diffraction datasets for structure determination. The crystal structure of TBEV NS5 MTase was solved at 1.9 Å resolution in space group P1 by molecular replacement, with two MTase molecules in each crystallographic asymmetric unit adopting highly consistent conformation (root-mean-square deviation (RMSD) value for all superimposable α-carbon atoms of 0.3 Å) (Table 1 and Supplementary Figure S1). Comparing with previously reported flavivirus MTase structures in isolation, the TBEV MTase adopts the same Rossmann fold with identical topology and very similar conformation, with corresponding RMSD values of 0.6–1.2 Å (chain A of the TBEV structure as the reference; 96–99% residue coverage) (Figure 1A). An S-adenosyl-L-methionine (SAM) molecule, the methyl donor of the methylation reaction, was bound in its binding pocket of each MTase.

The structure of TBEV NS5 RdRP was solved at 3.2 Å resolution in space group P212121 by molecular replacement, with ten RdRP molecules in each crystallographic asymmetric unit assembled in a dimer of pentamer fashion (Table 1 and Supplementary Figure S2, A–D), even though both the full-length NS5 and RdRP are monomeric in solution (Supplementary Figure S2E). Moreover, there is no conserved inter-molecular interface documented in available crystal forms of full-length and RdRP-only NS5 crystal structures. The packing among these dimer of pentamers resulted in spacious solvent channels in the crystal lattice, likely accounting for the poor-diffracting feature of the crystal (Supplementary Figure S2, A–D). The RMSD values are 0.4–0.5 Å (98–100% residue coverage).
Figure 1. Global views of TBEV NS5 MTase and RdRP structures. (A) Superimposition of the MTases of flaviviruses (left) and stereo-pair view of TBEV MTase in complex with SAM (right). Coloring scheme: left – TBEV MTase in cyan, MEAV MTase in dark turquoise, the MODV and YOKV MTase in red, and mosquito-flavivirus MTases in gray; right – subdomain I in violet, subdomain II in cyan, subdomain III in bright orange, SAM in green and zinc ions (spheres) in brown. The 3500 K composite SA-omit 2Fo – Fc electron density map of SAM was overlaid (contoured at 1.0 σ). (B) Superimposition of flavivirus RdRPs as cartoon representation. Coloring scheme: TBEV RdRP in dark orange, and mosquito-flavivirus (DENV, WNV) RdRPs in gray. (C) Global views of TBEV RdRP crystal structure. Structure of TBEV RdRP construct shown in orientations viewing into the front channel (left) and NTP entry channel (right), respectively. Coloring scheme: linker in red, RdRP palm in gray, N-terminal extension (NE) in violet, index finger in green, middle finger in orange, ring finger in yellow, pinky finger in light red, thumb in slate, priming element in purple, region B–C in light pink, motif C signature sequence SGDD in magenta. The numbers defining the residue ranges of each element are shown. (D) A color-coded bar of functional or structural elements of TBEV RdRP. The coloring scheme is the same as in panel C, and the unresolved regions in pinky/ring and at both termini are indicated by dark gray bars.

among these chains and 0.9–1.4 Å (74–93% residue coverage) among the RdRP-only flavivirus NS5 structures for all superimposable α-carbon atoms (both using chain A of the TBEV structure as the reference) (Figure 1B and Supplementary Figure S2F). The TBEV structure adopts the encircled human right hand architecture that is RdRP-unique, with palm, fingers, and thumb domains surrounding the polymerase active site (Figure 1B–D). Consistent with available flavivirus RdRP-only structures, part of the ring and pinky finger subdomains (residues 458–475 in ring and 408–416 in pinky for the TBEV structure) is disordered (20,41). The dynamics in the ring/pinky fingers, and in some cases also in index finger are also observed in full-length NS5 structures (18,39). To date, two types of full-length NS5 global conformations have been identified by crystallography. The conformation represented by the JEV NS5 structure typically has a completely folded fingers domain stabilized by intra-molecular MTase-RdRP interactions fea-
Figure 2. The structure and sequence diversity of region B-C of the *Flaviviridae* NS5 RdRs. (A) Structural comparison of the motifs B-C region of *Flaviviridae* NS5s. In the first row, TBEV is from tick-borne flaviviruses, the other four are from mosquito-borne flaviviruses. In the second row, TBEV and JEV are from *Flavivirus* genus, and HCV and CSFV represent other two genera of the *Flaviviridae* family. Coloring scheme: motif B in yellow, the region B-C in orange, motif C in light green. Two absolutely conserved residues (the G in the SG sequence of motif B and the first D in XGDD of motif C), and the two end residues of the structurally most diverse part of region B-C are shown as spheres. The /H9251-helix in motif B is shown as a cylinder. The 3500 K composite SA-omit 2Fo – Fc electron density map of TBEV region B-C was overlaid (contoured at 1.0). (B) Structure-assisting sequence alignment of the motifs B-C region of *Flavivirus* genus NS5s. Conserved active site residues are highlighted in red. Depending on the host range, viruses in the *Flavivirus* genus are divided into the four groups: the mosquito-borne group, the tick-borne group, the insect-specific group, and the vertebrate-specific group.

The conformational similarity between the TBEV RdRP and the mosquito-borne flavivirus RdRPs nevertheless supports the conservation of NS5 structure and conformation. However, whether the tick-borne flavivirus NS5 can indeed adopt the JEV- and DENV3-like global conformations requires further structural evidence.
The region between RdRP motifs B and C exhibits structure and sequence diversity and may have host adaptation-related features

While examining the structural difference between TBEV RdRP and other flavivirus RdRPs, the region between RdRP catalytic motifs B and C (hereinafter termed ‘region B-C’) drew our attention. Motifs B and C are among the most conserved regions of viral RdRPs. Forming a β-hairpin substructure, motif C contains an absolutely conserved aspartic acid residue (the first ‘D’ in the RdRP signature sequence XXDD) in the hairpin region that participates in catalytic divalent metal ion coordination. Motif B is consisted of an N-terminal loop region and a C-terminal helical region, and contains an absolutely conserved glycine residue in the loop (the ‘G’ in a highly conserved SG sequence) that plays an important role in NTP recognition (42,43). Region B-C starts in the central part of the long helix that harbors the motif B helical region and ends at the beginning of the N-terminal β-strand in motif C (Figure 2 and Supplementary Figure S3). This region resides at the bottom of the RdRP palm and is solvent accessible, and therefore has the potential to interact with viral and/or host factors or to mediate processes other than RNA synthesis. Compared to its neighboring motifs, region B-C exhibits relatively high structural variance and dynamics (Supplementary Figure S3, B–C). We previously carried out a structure and function survey of RdRP from positive-strand RNA viruses and found that the length of region B-C varies to a great extent, with a length difference greater than 60 residues between the longest and the shortest (44). In flaviviruses, the length of this region is about 40 residues and exhibits length and sequence diversity between different groups. The TBEV RdRP region B-C is about 2–3 residues shorter than its mosquito-borne flavivirus counterparts (Figure 2B). If compared with RdRP structures of known mosquito-borne flaviviruses, residues 632–640 of TBEV RdRP or 633–644 of JEV RdRP represent the structurally most diverse part of this region (Figure 2A, top panel). Comparing to HCV and CSFV NS5B proteins, two representative Flaviviridae RdRPs, region B-C of flavivirus RdRP is about 15–25 residues longer (Figure 2A, bottom panel).

Inter-species region B-C substitutions do not abrogate RdRP catalytic activities

In order to further explore the role of region B-C in flaviviruses, we designed a set of four inter-species substitution mutations in TBEV and JEV NS5s. The residue ranges chosen for substitution corresponding to the solvent exposed part of region B-C (TBEV NS5 632–651) or the structurally most different stretch (632–640), resulting in two pairs of mutations with long (JEV LS and TBEV LS) and short
(JEV SS and TBEV SS) substitutions, respectively (Figure 3). Corresponding TBEV and JEV NS5 mutant constructs were then compared with WT enzymes in both primer-dependent and dinucleotide-driven RdRP assays to assess whether substitutions in region B-C affect polymerase catalytic activities. In previous studies characterizing flavivirus NS5 RdRP, DENV2 NS5 exhibited manganese ion (Mn$^{2+}$)-independent activities in vitro, while JEV NS5 required Mn$^{2+}$ in both primer-dependent and dinucleotide-driven assays (38,39). It turned out that TBEV NS5 also needs Mn$^{2+}$ in our experimental settings. Using a 33-nucleotide (nt) RNA template (T33) and a 10-nt primer (P10) that can form an 8-base pair (bp) template-primer duplex, an A-U-A-U tetra-nucleotide was readily incorporated to make a 14-nt product (P14) within 5 min using WT TBEV and JEV enzymes, while a minor product 15-nt in length likely derived from a misincorporation event was also observed (Figure 4A–B; Supplementary Figure S4, lanes 2–6). Region B-C substitutions in either TBEV or JEV NS5 did not apparently affect the product profiles and formation kinetics (Figure 4B, compare lanes 2–4 with lanes 5–7 and 8–10 for TBEV constructs; compare lanes 22–24 with lanes 25–27 and 28–30 for JEV constructs). The dinucleotide-driven assays were typically used in various polymerase systems to mimic de novo synthesis (45–47), and a 30-nt template (T30) and a G–G dinucleotide with a 5′-phosphate (pGG) previously used in characterizing JEV, DENV and CSFV RdRPs were used in this study (Figure 4C). When ATP and UTP were used as the only NTP substrates, the formation of the 9 nt product for TBEV WT NS5 was slower than that of JEV WT enzyme (Figure 4D, compare lanes 12–14 with 32–34). The 10 nt product was observed in both enzyme systems, possibly derived from a G:U misincorporation event (Supplementary Figure S4, lanes 7–9). For substitutions using the TBEV RdRP backbone, no obvious effect on catalysis was observed (Figure 4D, compare lanes 12–14 with lanes 15–17 and 18–20). By contrast, substitutions in the JEV RdRP backbone resulted in slower formation of the 9–10 nt products (Figure 4D, compare lanes 32–34 with lanes 35–37 and 38–40). Taken together, the maintenance of RdRP
Figure 5. The inter-species region B-C substitution abolished virus proliferation. (A) Schematic diagram of WT infectious clones and the corresponding TBEV_SS, JEV_SS and JEV_LS mutant clones. For all three substitution mutants, the design of substitution is identical to those in Figure 3. (B) IFA analysis of WT and TBEV_SS, JEV_SS and JEV_LS mutants. Equal amounts of genomic RNAs of JEV WT, JEV_SS and JEV_LS mutants were transfected into BHK-21 and C6/36 cells, respectively. IFA was performed using JEV E (4G2) and TBEV NS5 antibodies to detect the viral protein synthesis at the indicated timepoints. (C) The viral RNA quantification of TBEV. The total mRNA of BHK-21 cell was extracted, and TBEV mRNA was detected at 1, 2 and 3 d.p.i. via RT-qPCR. (D) The western blotting assay detecting TBEV NS5 in the infected BHK-21 cells. GAPDH (glyceraldehyde-3-phosphate dehydrogenase) was used as a loading control.

activities after region B-C substitutions in both TBEV and JEV systems suggests that region B-C may not play essential roles in RdRP catalysis.

Inter-virus region B-C substitution abolishes TBEV and JEV proliferation but supports low-level RNA replication in a TBEV replicon

In order to assess the role of region B-C in flavivirus proliferation, SS substitution was introduced into a TBEV infectious clone, while the corresponding LS and SS substitutions were introduced into a JEV infectious clone (Figure 5A) (48–50). For the virus recovery of WT and mutant TBEV, as described previously (49,50), equal amounts of infectious clone plasmids were transfected into BHK-21 cells to assess the effect brought by the substitution. Three days post transfection, the cells transfected with the WT infectious clone plasmids exhibited obvious cytopathic effect, and supernatant samples (for both WT and mutant) were then prepared and used to infect naive BHK-21 cells (with a virus titer of $7 \times 10^6$ plaque forming units (pfu)/ml determined by a plaque assay) (Supplementary Figure S5). Viral NS5 protein production level was monitored by an immunofluorescence assay (IFA). IFA positive cells were not detected for 3 days post infection for the mutant clone, while obvious NS5 production was evident for the WT clone (Fig-
ure 5B, top three rows). For the WT and mutant JEV clones, equal amounts of viral RNAs were transfected into BHK-21 or C6/36 cells to assess the effect brought by the mutations in both mammalian and mosquito host cells. Viral envelope (E) protein production level was monitored by IFA. In either host cell systems, IFA positive cells were not detected for both LS and SS substitution clones after transfection, while continuous production of viral E protein was observed for the WT clone (Figure 5B, bottom two rows). The TBEV IFA data were further validated by comparing the amount of viral RNA copies and antibody-based NS5 production levels, and the SS mutant virus failed to produce viral RNAs and NS5 proteins (Figure 5C–D). These data together suggest that region B-C plays critical roles in virus proliferation for both virus species. Although the LS/SS mutations may change the local RNA structure, it is less likely for these nucleotide mutations to abolish virus proliferation by disrupting conserved long-range interactions essential to flavivirus replication mainly involving conserved structural elements in the untranslated regions (51,52).

To further explore the functional relevance of region B-C, a TBEV replicon bearing the SS substitution and a luciferase reporter gene was constructed, and the luciferase activity and RNA copy numbers were used to assess the replicon-derived RNA replication in BHK-21 cells transfected with the replicon RNAs (Figure 6). Interestingly, the SS mutant replicon was able to support low level replication as indicated by the luciferase activities and RNA copy numbers if compared with WT replicon (about 1 order of magnitude lower than the WT), but it clearly replicated better than a mutant replicon bearing double D-to-A mutations in both mammalian and mosquito host cells. Vi-
Figure 7. A diversity survey of region B-C region in viral RdRPs. (A) Structural comparison of the motifs B-C region in RNA virus polymerases. Each structure represents an RNA virus family. Coloring scheme: motif B in yellow, region B-C in orange, motif C in light green. Two absolutely conserved residues (the G in the SG sequence of motif B and the first D in XGDD of motif C) are shown as spheres. The α-helix in motif B is shown as a cylinder. ss (+) RNA, ss (−) RNA, ds RNA are used as abbreviations for positive-strand, negative-strand, and double-strand RNA viruses, respectively. (B) Structure-based sequence alignment of RdRP motifs B and C, with region B-C sequences listed (not aligned) for comparison. Two absolutely conserved residues are highlighted in red.
regions and co-foiling with other viral proteins (68,69), the RdRP catalytic module structure is relatively conserved. In particular, the spatial arrangement of motif A-G is consistent for all RdRPs despite low sequence conservation level in the scope of all RNA viruses (54,70). Typically, the seven motifs appear in the order of G-F-A-B-C-D-E in RdRP primary structure, except for the Permutatetraviridae and Birnaviridae RdRPs that have a different order of G-F-C-A-B-D-E (57,71). Among them, motifs A, B, C, and F directly interact with the NTP substrate and contain highly conserved residues (42,72). In contrast, motifs D, E and G located at the periphery of the active site mainly play structural roles and are less conserved in sequence (70,73). Hence the residue spacing among motifs F-A-B-C can be readily surveyed in majority of RdRPs by locating the highly conserved residues. While the residue spacing for motifs F/A and A/B is relatively consistent, and is about 40–60 residues for both regions, the spacing between motifs B and C varies to a much greater extent (44,55). The shortest spacing is smaller than 20-residue as seen in RdRPs from Tymoviridae, Tobagoviridae, Alvernaviridae etc., while the longest is larger than 70-residue in Mononiviridae (belonging to Nidovirales) RdRPs in a survey of RdRPs from positive-strand RNA viruses (44).

The observations of large distance variation between RdRP motifs B and C echo the aforementioned structure and length variations among flavivirus and Flaviviridae RdRP region B-C, and invite us to further survey the corresponding region of other representative RdRP structures. For other positive-strand RNA viruses, region B-C lengths of RdRPs from Picornaviridae, Caliciviridae and Leviviridae are relatively short, while those from Coronaviridae are the longest and forming a helix bundle (Figure 7A, top panel; Figure 7B). As revealed in the SARS-CoV-2 RdRP-nsp7-nsp8 complex structure, part of region B-C interacts with the NiRAN domain hairpin, while the rest of it is solvent accessible (65). For double stranded RNA viruses, the Reoviridae and Picorniviridae RdRP region B-Cs are short and comparable to the Picornaviridae counterparts, while that from the Cystoviridae is longer and similar to the flavivirus region B-C both in length and structure (Figure 7A, middle panel; Figure 7B). For negative-strand RNA viruses including Pneumoviridae, Rhabdoviridae, Orthomyxoviridae, and Peribunyaviridae, the lengths of RdRP region B-C are relatively short, but the spatial placement (orientation) of this region relative to RdRP active site varies to some extent (Figure 7A, bottom panel; Figure 7B).

Taken together, the length and structure diversity of region B-C and its solvent accessible feature suggest that it may have become a regulatory hotspot in RdRP evolution history. Inter-species substitution retained RdRP catalytic activities but did not support virus proliferation further support the possibility of its involvement in other viral/host events. Through identifying relatively subtle differences of region B-Cs between the tick-borne and mosquito-borne flavivirus RdRPs and generalizing its high-level of diversity in the context of viral RdRPs, the current study pointed out this probable host adaptation-related region within the most conserved enzyme module of RNA viruses. However, the specific role of this region and the precise mechanism underneath, either in flavivirus or in others, require further investigations including discoveries of region B-C interactions with viral and host elements.

DATA AVAILABILITY

The atomic coordinates and structure factors for the reported crystal structures of the TBEV NS5 MTase and RdRP have been deposited in the Protein Data Bank under accession numbers 7D6M and 7D6N, respectively.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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