Structure of the St. Louis encephalitis virus postfusion envelope trimer

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St. Louis encephalitis virus (SLEV) is a mosquito-borne flavivirus responsible for several human encephalitis outbreaks over the last 80 years. Mature flavivirus virions are coated with dimeric envelope (E) proteins that mediate attachment and fusion with host cells. E is a class II fusion protein, the hallmark of which is a distinct dimer-to-trimer rearrangement that occurs upon endosomal acidification and insertion of hydrophobic fusion peptides into the endosomal membrane. Herein, we report the crystal structure of SLEV E in the posfusion trimer conformation. The structure revealed specific features that differentiate SLEV E from trimers of related flavi- and alphaviruses. SLEV E fusion loops have distinct intermediate spacing such that they are positioned further apart than previously observed in flaviviruses but closer together than Semliki Forest virus, an alphavirus. Domains II and III (DII and DIII) of SLEV E also adopt different angles relative to DI, which suggests that the DI-DII joint may accommodate spheroidal motions. However, trimer interfaces are well conserved among flaviviruses, so it is likely the differences observed represent structural features specific to SLEV function. Analysis of surface potentials revealed a basic platform underneath flavivirus fusion loops that may interact with the anionic lipid head groups found in membranes. Taken together, these results highlight variations in E structure and assembly that may direct virus-specific interactions with host determinants to influence pathogenesis.

St. Louis encephalitis virus (SLEV) was first discovered as the agent responsible for over 1,000 cases of encephalitis during a 1933 summer outbreak in St. Louis, MO (1, 2). Additional epidemics have occurred from 1964 to 2006 in the Americas, ranging from the United States (3–5) to Argentina and Brazil (6–8). However, reported infections have declined significantly in the United States since West Nile virus (WNV) emerged in 1999 (9), suggesting WNV has since outcompeted SLEV in North America. SLEV naturally cycles between Culex, Aedes, Anopheles, and Theobaldia genera (10, 11) mosquitoes and passerine birds, but humans may also become infected as “dead end” hosts through mosquito bites.

SLEV is a member of the Flavivirus genus of the Flaviviridae family. Other flaviviruses known to infect humans include dengue virus (DENV), Japanese encephalitis virus (JEV), and tick-borne encephalitis virus (TBEV). Each flavivirus is classified into a serocomplex with its own distinct tropism and pathogenesis, based on cross-neutralization tests (12, 13). More specifically, when polysera harvested from an animal infected with one virus neutralizes a different virus in vitro, the two are said to belong to the same serocomplex. SLEV, JEV, and WNV are members of the JEV serocomplex, while DENV and TBEV each represent prototypical members of other prominent serocomplexes. JEV serocomplex viruses are transmitted by mosquitoes and often cause asymptomatic infections, but severe cases can result in encephalitis, fever, and meningoencephalitis (14). TBEV, Langat virus, and Powassan virus make up the TBEV serocomplex and are transmitted by hard ticks. These viruses cause similar illness to that of the JEV serocomplex but spread to secondary tissue more slowly and are more frequently associated with severe symptoms (15). The DENV serocomplex contains 4 serotypes (DENV1 to -4) of mosquito-borne viruses. Infection with any DENV serotype can cause headaches, muscle pain, skin rash, and hemorrhagic fever.

The flavivirus virion is enveloped, and its surface is decorated with 180 transmembrane envelope (E) proteins (16, 17) that mediate cellular attachment and fusion with host membranes. Flavivirus E, along with the alphavirus E1 protein, represent class II viral fusion proteins based on their activation mechanisms and three-dimensional structures (18–20). Features that define class II proteins are a requisite dimer-to-trimer rearrangement prior to fusion and a distinct β-strand-rich, 3-domain architecture (18–20). The SLEV E ectodomain is 68% identical to serocomplex-related JEV E but only 46% and 40% identical to those of DENV2 and TBEV, viruses from different serocomplexes. Alphavirus E1, however, shares no appreciable sequence identity to SLEV E or any other flavivirus E protein. Despite differences in their amino acid sequences and arrangements on the viral particle, the structures of E and E1 are remarkably similar (16, 21–25). The conservation of three domains has been well documented in crystal structures of flavivirus E ectodomains from TBEV (24), DENV (26), JEV (27), and WNV (28, 29), as well as alphavirus E1 from Semliki Forest virus (SFV) (25) and Chikungunya virus (CHIKV) (30). Domain I (DI) is a β-barrel, DII is formed from two discontinuous loops that protrude from DI, and DIII is an Ig-like sandwich positioned on the opposite side of DI from DII. Antibodies that neutralize SLEV (31, 32), WNV (33–39), DENV (40–43), and JEV (44, 45) have been mapped to epitopes on all three domains. These antibodies can inhibit both attachment to cells and fusion in the endosome (46–48).
of the hydrophobic fusion loop at the dimer interface to prevent premature insertion into host membranes. Upon internalization, the acidic pH of the endosome disrupts the E dimers and transforms them into a fusogenic state (49, 50). E inserts its fusion loops into the endosomal membrane and collapses into a compact, trimeric conformation that forms when DIII swings in an arc to pack against DI beneath DII (21, 23, 49, 51–53). Alphavirus E1 forms heterodimers with chaperone protein E2 but dissociates and undergoes a similar conformational change when acidified in the late endosome (54). In either case, this transition is believed to provide the energy to fuse viral and host membranes, releasing the nucleocapsid into the cytosol.

We previously established that prefusion E proteins of the JEV serocomplex have distinct biophysical characteristics, such as a smaller dimer interface and diminished dimerization propensity (27, 29). We therefore determined the crystal structure of acidified SLEV E to examine the structure and assembly of the trimeric, postfusion conformation. SLEV E served as a representative of the JEV serocomplex for comparative analysis with class II fusion proteins from DENV and TBEV serocomplexes and the distantly related alphavirus SFV. SLEV E did not crystallize in the prefusion conformation, so we utilized the structure of serocomplex–related JEV E as a model to evaluate probable differences between prefusion and postfusion SLEV E. Our analyses revealed that postfusion envelopes possess a series of well-conserved trimer contacts yet exhibit substantial differences in domain orientation, fusion loop spacing, and electrostatic surface potential. These biophysical properties may influence pathogenesis by modulating interactions with environmental factors, such as the specific fusion pH or host membrane composition.

**MATERIALS AND METHODS**

**Cloning, expression, and purification of SLEV E.** Residues 1 to 407 of the SLEV E ectodomain (strain MS1-7) were cloned into the pET21a(+) expression vector. The vector was transformed into *Escherichia coli* BL21-CodonPlus(DE3)RIL cells. An individual colony was inoculated to inoculate four 10-mL starter cultures that were transferred to 4 liters of methionine-deficient medium (Athena Enzyme Systems). Cultures were grown at 37°C until reaching an optical density (OD) of 0.6. Upon reaching this OD, cultures were supplemented with 0.5 g feedback inhibition stock (0.1 g L-lysine, 0.1 g L-threonine, 0.1 g L-phenylalanine, 0.05 g L-leucine, 0.05 g L-isooleucine, 0.05 g L-valine, and 0.05 g L-selenomethionine) to prevent synthesis of methionine and initiate incorporation of selenomethionine (protocol adapted from that described by Stols et al. [55]). At 15 min after addition of this stock, 1 ml of 1 M isopropyl-β-D-thiogalactopyranoside (IPTG) was added to each 1-L culture to induce protein expression. After 6 h, cultures were harvested and cells were pelleted. Cells were solubilized in solution buffer (50 mM Tris [pH 8.0], 25% sucrose, 10 mM dithiothreitol [DTT]), and an equal amount of lysis buffer (50 mM Tris [pH 8.0], 1% Triton X-100, 100 mM NaCl, 10 mM DTT) was added. Lysozyme was added to a final concentration of 0.8 mg/ml, and the mixture was sonicated to disrupt cellular membranes. This solution was then centrifuged at 10,000 rpm to pellet inclusion bodies. The inclusion bodies were washed three times with wash buffer (50 mM Tris [pH 8.0], 0.5% Triton X-100, 100 mM NaCl, 1 mM DTT) and sonicated after each wash. Finally, the mixtures were washed in buffer without Triton X-100 and then solubilized in 6 M guanidine-HCl, 10 mM Tris (pH 8.0), and 20 mM β-mercaptoethanol. Aliquots of this solution were added dropwise to a reservoir of refolding buffer containing 400 mM non-detergent sulfobetaine 201 (NDSB-201), 100 mM Tris (pH 8.0), 0.5 mM oxidized glutathione, and 5 mM reduced glutathione and allowed to fold overnight. Protein was then concentrated using an Amicon 400 concentrator with 30-kDa cutoff membrane and purified by size exclusion chromatography (SEC) and anion-exchange chromatography.

**Crystallization.** Purified, soluble SLEV E was dialyzed into a buffer containing 0.05 M acetate (pH 5.5) and 10 mM sodium chloride and crystallized by hanging drop vapor diffusion at 20°C. Each drop was a mixture of 0.5 μl of protein solution at a 5-mg/ml concentration and 0.5 μl of mother liquor containing 0.1 M acetate (pH 5.5), 3% polyethylene glycol 8000 (PEG 8000), and 2% ethylene glycol. Diffraction quality crystals grew in 3 to 7 days. Crystals were rapidly dropped through a drop of 25% ethylene glycol for cryoprotection prior to cooling in liquid nitrogen. Data were collected at the APS (advanced photon source) 19-ID beamline and processed, integrated, and scaled using HKL2000 (36). SLEV E crystallized in space group I23, with unit cell dimensions of a = b = c = 177.5 Å and α = β = γ = 90°.

**Structure determination.** A model of SLEV E in the postfusion conformation was utilized to solve the structure by molecular replacement (MR). This model was generated with the Phyre2 server (57) by threading the SLEV E sequence onto DI and DII of DENV2 E in the postfusion trimer conformation (PDB ID 1OKB) and onto DII of JEV E (PDB ID 3P54). MR was performed with the PHENIX program (58). Refinement was carried out by using a variety of approaches. PHENIX refine was used to perform rigid body refinement of the three domains, followed by positional refinement incorporating secondary structure restraints, simulated annealing, and TLS (translation liberation screw) refinement (59) using three groups and B-factor refinement. The resultant partial model was used to carry out a combination of MR and selenomethionine single anomalous dispersion (MRSAD) phasing in the PHENIX Autosol program (58, 60). The improved maps generated by the incorporation of the experimental phase allowed for model building in Coot (61) with the assistance of B-factor sharpening using a value of 15Å. Additional rounds of positional and B-factor refinement were performed using the jelly body function of refmac5 (62) or with PHENIX refine (58). The final structure had an Rwork of 22.2% and Rfree of 26.7% and contained ectodomain residues 2 to 145 and 163 to 404 of the 407 total encompassed by the model.

**Structural analysis.** Anomalous difference peaks were generated with PHENIX Autosol (58) and the CCP4 Fast Fourier transform (FFT) algorithm (63). First, SAD phasing was carried out in Autosol in the absence of a partial model to avoid model bias. Next, this solution and the SLEV E model were used as inputs for the CCP4 FFT to generate a map for use in visualization software. The resultant density was visualized in Pymol (64) at a contour level of 4.5σ. Interatomic distances between fusion loops were measured in Pymol. The PISA Web server (65) was utilized to determine the buried surface area and contacts between E protomers. ClustalW was used to align flavivirus E sequences (66). To determine angles between DI and DII or DIII, entire E proteins from DENV1, DENV2, TBEV, and SFV were superimposed onto SLEV DI by secondary structure matching in Coot (61). Next, artificial models of each E protein were generated by superimposing DI, DII, and DIII individually onto SLEV E to create a reference structure. The angles between DI and DII or DI and DIII of the native structure versus the reference structure were then calculated with Dymond (63, 67). Electrostatic surface potentials were determined by using PDB2PQR (68, 69) to generate PQR files with protonation states assigned at pH 6.0 for use in the APBS (70) plugin for Pymol and visualized over a range of −3 kT/e to +3kT/e.

**Protein structure accession number.** The coordinates for the protein structure have been deposited in the Protein Data Bank and assigned PDB ID number 4FG0.

**RESULTS**

**SLEV E structure.** A selenomethionine (SeMet)–substituted SLEV E ectodomain from strain MS1-7 lacking the C-terminal stem region was overexpressed as bacterial inclusion bodies. These inclusion bodies were solubilized and oxidatively refolded to allow for proper disulfide formation. This strategy was employed be-
cause initial crystals of SLEV E diffracted poorly, and so the experimental phase information gained through this incorporation could be used to validate the structure through the identification of anomalous SeMet peaks in the electron density maps. Purified SLEV E crystallized in cubic space group I23, and phase was determined using a combination of MR and SAD data. There was one E monomer per asymmetric unit, but application of cubic symmetry generated the postfusion trimer (Fig. 1A). The structure was refined at 3.9-Å resolution with an $R_{work}$ of 22.2% and $R_{free}$ of 26.7% (Table 1), and it included ectodomain residues 2 to 145 and 163 to 404. Residues 146 to 162 of the loop between the Eo and Fo strands were disordered, as has been observed in other postfusion E structures (23, 51, 52).

SLEV E adopted the three-domain architecture characteristic
of class II viral fusion proteins (Fig. 1A). DI is a β-barrel composed of 8 strands, and DII is formed from two extended loops that protrude from DI. DII contains a long β-sheet made up of strands in both portions of this discontinuous domain that supports the fusion loop. At the opposite end of DI is a flexible linker that connects to DIII, a 7-stranded β-sandwich similar to the classical Ig fold. Since the structure was determined at low resolution, anomalous peaks generated through SAD phasing in the absence of a model were used for validation. Each of the 7 SLEV E sel enomethionine residues localized to the density at these peaks (Fig. 1B), confirming the sequence register and accuracy of the enomethionine residues localized to the density at these peaks.

Differences in JEV serocomplex E structures. Crystal structures of JEV serocomplex E proteins from SLEV, WNV, and JEV provided a detailed structural perspective of the viral life cycle. While it would have been ideal to compare the pre- and postfusion conformations from SLEV E, extensive trials only yielded diffracting crystals of the postfusion trimer. In the absence of such a structure, we believe monomeric and prefusion E proteins from related serocomplex members WNV and JEV provided a suitable basis for comparison. WNV E crystallized in an unusual monomerization that may represent a structural transition that occurs during maturation or acidification. The WNV E proteins assembled in a perpendicular lattice with fusion loops buried in the same pocket they occupy in the mature dimer (29). JEV E formed the antiparallel prefusion dimer observed in cryo-EM structures of mature virions, with fusion loops buried in a cavity formed by DI and DIII of the opposing subunit (27). SLEV E, on the other hand, crystallized as a postfusion trimer (Fig. 2A). In this conformation, DIII swings 63° toward DI relative to its equivalent position in the JEV E dimer. This reorientation causes it to interact with its parent DI, instead of DII of a neighboring protomer.

Several loops that protrude from the DI β-barrel toward DIII have specific conformations that accommodate the pre- or postfusion DI-DIII linker. The JEV prefusion linker lies between the Cα-Dα and Gα-Hα strands (labeled as in Fig. 2B) and emerges from DI as an essentially linear peptide. In the postfusion structure, this peptide kinks at residues D291/K292 and threads back toward DI between the Aα-Bα and Cα-Dα loops (Fig. 2B). The position of the prefusion linker results in a shift of the Gα-Hα loop outward from the DI core relative to the postfusion structure. However, the postfusion DI Aα-Bα and Cα-Dα loops are splayed apart to permit this alternate trajectory.

Trimer packing and assembly. Comparison of trimer and dimer contacts from a variety of flavivirus E structures revealed two unique interfaces, with the exception of a few overlapping residues near the fusion loop (Fig. 3B). Thus, the chemical changes induced by the acidic pH of the endosome enable the formation of entirely new interfaces. SLEV E trimers are stabilized by a series of interactions involving all three domains. The lateral edge of DI from one protomer packs into a groove formed by DI and DIII of its neighbor to secure the base of the trimer (Fig. 3A). Additionally, the edge of DII contacts residues of an adjacent DII near the DI-Dll joint and beneath the fusion loop. SLEV trimers were generated by application of 3-fold crystallographic symmetry, so that the two interfaces were identical on each protomer.

We previously reported that JEV E possesses substantially less buried surface area than other dimeric E structures and does not favor dimerization at prefusion (neutral) pH, as evaluated by multilevel light scattering (27). SLEV E and WNV E also did not dimerize under these conditions, suggesting that the icosahedral assembly of JEV serocomplex dimers is stabilized by lateral contacts present in mature virions. We therefore compared contact residues and the buried surface area of SLEV E relative to DENV1/2 and TBEV to investigate serocomplex-specific features of postfusion trimer structure and stability. While JEV E dimers buried ~50% less surface area than DENV or TBEV E, the extensive SLEV E trimer interface buried 2,100 Å² ~10% more than DENV1/2 E and the same amount as TBEV E or SVF E1 (Table 2).

Class II trimer relative domain orientations. Class II envelope proteins are comprised of three rigid domains linked by two joints (24, 30, 71, 72). These flexible regions connect the central DI to DII on one side and DIII on the other. Conformational changes borne out of twisting or bending of domains about these residues are vital to many stages of the flavivirus life cycle. SLEV DI was used as a reference for structural alignments to compare relative domain orientations in postfusion envelope structures from TBEV, DENV1, DENV2, and SFV (Fig. 4A). The angles relating DI-DII and DI-DIII varied considerably among the trimeric structures. DII from DENV1 and DENV2 adopted similar conformations, shifted ~11° relative to SLEV DII (Table 2). TBEV and SFV exhibited ~9 to 10° differences in the DI-DII angle relative to SLEV but are tilted in a different direction than those of DENV1/2 (Fig. 4A; Table 2). When viewing the fusion loops from a top-down perspective, DI does not appear to have crystallized in positions along a single trajectory but instead pivots about DI (Fig. 4B). These variable DI angles suggest that the DI-DII joint operates like a ball and socket rather than a hinge.

The relative positions of DIII vary over an even larger range than those of DII. DENV1 and DENV2 DIII are tilted 6.5° and 2.5°

### Table 1: Data collection and refinement statistics

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<tr>
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<tr>
<td>Cell dimensions</td>
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<td>a = b = c = 177.5 Å,</td>
<td></td>
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<tr>
<td>α = β = γ = 90.0°</td>
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<td>Redundancy</td>
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<td>Rmerge (%)</td>
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<td>R-free (%)</td>
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<td>Rotamer outliers (%)</td>
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*Values in parentheses are for data in the highest-resolution shell. Statistics are as defined in the Phenix program.*
in comparison to that of SLEV, while TBEV DIII is rotated 16.5° in this same direction about DI (Fig. 4A). SFV DIII was rotated 65° relative to SLEV DIII, representing a dramatic difference in the conformation of postfusion alphavirus versus flavivirus envelopes. This twisting of DIII alters the overall trimer geometry, interaction surface with DI, and potentially the direction the stem-loop projects from its C terminus.

**Fusion loop separation.** The different DI-DII angles and trimer assemblies observed among flavi- and alphavirus envelopes alter the distance between fusion loops. SFV E1 fusion peptides are positioned 47 Å apart and adopt an uncoiled conformation structurally distinct from those of the flaviviruses (Fig. 5). A length of 26 Å separates the loops of DENV1, DENV2, and TBEV E, while SLEV E fusion loops are situated at an intermediate distance of 35 Å (Fig. 5; Table 2). Trimer interfaces are well conserved in these E proteins, so this variable fusion loop spacing can be attributed to changes in DII tertiary structure and pivot positions about DI. It should be noted that the recombinant SLEV E is selenomethionine substituted, but any perturbations due to this incorporation will be local and unlikely to result in changes in domain orientation.

**Trimer electrostatics.** To investigate electrostatic features of E trimers that may contribute to structural changes or interactions with host determinants, we visualized the surfaces on postfusion E or E1 from DENV1, DENV2, TBEV, SLEV, and SFV. Protonation states were assigned at pH 6.0 to estimate the acidic environment of the endosome. In the flavivirus trimers, uncharged crowns of fusion loops lie atop a basically charged platform (Fig. 6A to D). Given that the fusion loops project into the hydrophobic portion of the bilayer, it seems reasonable to propose that the basic platform stabilizes the membrane-trimer complex by interacting with
negatively charged lipid head groups. An acidic ridge of DII was also visible between SLEV E protomers. Repulsive forces exerted at this interface could serve to distort these domains (Fig. 6D) and position the fusion loops further apart than those of DENV1/2 or TBEV. An acidic patch is also present on the interior face of SFV fusion loops facing the central axis (Fig. 6E). These regions may repulse one another to splay fusion loops apart in the flower-like arrangement observed in the SFV trimer (21).

**DISCUSSION**

The crystal structure of SLEV E revealed distinct biochemical features that may influence the interaction of JEV serocomplex vi-

<table>
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<th>Virus (trimer)</th>
<th>Distance between fusion loops (Å)</th>
<th>DI-DII angle (*) relative to SLEV E</th>
<th>DI-DIII angle (*) relative to SLEV E</th>
<th>Buried surface area (Å²)*a</th>
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<td>SFV (1RER)</td>
<td>47</td>
<td>9.9</td>
<td>65.2</td>
<td>2,139*</td>
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</table>

*a*, the buried surface area was averaged over each trimer subunit.

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**FIG 3** Conservation of flavivirus E trimer interfaces. (A) The left panel displays trimer interface 1, formed by two E protomers. Residues contacted by the cartoon representation are highlighted on the surface-rendered protein. Green residues represent contacts exclusive to the surface-rendered protein, while cyan residues are contacts in both molecules at the interface. The third E protomer that does not make contacts at the highlighted interface is shown as a transparent mesh envelope. (B) The right panel displays trimer interface 2. Contacts in both molecules in this panel are also cyan, while residues exclusive to the surface-rendered protein are shown in gray. The protomer that does not make contacts at this interface is also shown in mesh representation. (C) A ClustalW sequence alignment of SLEV, DENV1, DENV2, and TBEV E. SLEV E domains I, II, and III are shown in red, yellow, and blue, respectively. The SLEV E secondary structure is displayed above the sequence with a straight line for coils/loops, arrows for strands, and cylinders for helices. Contact residues are highlighted in the same color scheme described for panels A and B. For JEV, DENV2, and TBEV E proteins with dimeric structures available, the dimer contacts are underlined in black.
ruses with host determinants, and therefore pathogenesis. In our analyses, SLEV E served as a basis for a structural comparison between all postfusion class II structures. Domains II and III of these envelope proteins are able to accommodate a range of orientations while remaining part of the trimeric assembly. Additionally, orientation of these domains contributed to variations of distances between the triangular crowns of fusion loops. Fusion loops of SLEV E were separated by an intermediate length compared to the tight spacing of DENV1, DENV2, and TBEV E or the wide set loops of SFV. Models of flavivirus fusion propose that the stem region truncated in recombinant E lies between trimers and splays them apart so that they reach an alphavirus-like conformation (23,51, 73, 74). Recombinant SLEV E does not contain this stem, but it crystallized with wider spacing than other E trimers. This fusion loop separation could thus represent a characteristic feature of SLEV E or the JEV serocomplex.

Despite variations in relative domain positions, trimeric SLEV, DENV1, DENV2, and TBEV E proteins interact through at a series of structurally conserved positions (Fig. 3B). Prefusion dimers also assemble by forming contacts at conserved positions, but the dimer interface is generally smaller and appears to vary over a larger range. Also, SLEV E was previously found to dimerize less readily than DENV2 E (27), an observation that suggests the forces governing dimer and trimer assembly are not necessarily correlated. Trimer interfaces have similar buried surface area values, from 1,850 to 2,100 Å², while dimers buried at 840 to 1,930 Å². This wide range of calculated dimer surface area values may be explained by the presence of additional lateral interactions present outside the dimer interface in mature virions. On the other hand, the more conserved trimer interfaces may be necessary to rigidify E to allow for efficient fusion.

Ultimately, the functional reason fusion loops are exposed atop the E trimer is to drive interactions with the host lipid bilayer and merge it with that of the virion. This role led to the hypothesis that lipids or detergents would be required for trimer formation, and so the initial crystallization of postfusion E proteins from DENV2 (23) and TBEV E (51,75) was carried out by purifying trimeric E complexes in detergents to simulate their membrane environment. However, our SLEV E (pH 5.5) structure and a previous structure of DENV1 E (pH 6.5) (52) have now been crystallized in the postfusion state without detergent, indicating that high protein concentration and an acidic environment are sufficient for trimerization. Interestingly, crystals grown in the absence of detergent (SLEV E,, 3.9 Å; DENV1 E, 3.5Å) have so far diffracted to lower resolutions than those crystallized in the presence of detergent (DENV2 E, 2.0 Å; TBEV E, 2.7 Å). Since weak diffraction is often a result of molecular motion or vibration, it is possible that detergents enhance trimer stability but are not an absolute requirement.

Biochemical features of class II envelope proteins may be finely
tuned for interaction with environmental conditions, such as specific lipid compositions of membranes or pH values. Such interactions can direct viruses to certain host cells or cellular compartments. For example, alphavirus SFV requires cholesterol and sphingolipids for efficient fusion (76, 77), while fusion of flaviviruses DENV2 and DENV4 is dependent upon anionic lipids bis-(monoacylglycero)phosphate and phosphatidylserine, which are found only in the late endosome (78). Analysis of trimer surface

FIG 5 Fusion loop separation. Fusion loops are shown in green with side chain sticks, and the conserved glycine used as a reference point for distance measurement across loops is shown in magenta (top row). Postfusion trimers from TBEV, DENV2, SLEV, and SFV are displayed in cartoon representation; DENV1 loops were equivalently spaced to DENV2 but are not shown. DI, DII, DIII, and the fusion loop of one E subunit are shown in red, yellow, blue, and green, respectively (bottom row). The second and third protomers are shown in gray and wheat.

FIG 6 Trimer electrostatics. TBEV, DENV1, DENV2, SLEV (E) and SFV (E1) envelope surface representations are colored based on surface potential, with red as negative potential and blue as positive. The top panel is a top-down view of the fusion loops, and the bottom panel is a side view. SLEV and SFV have regions of negative charge that may be relevant to trimer structure or assembly (circled in black). The basic platform beneath flavivirus fusion loops is also labeled in panel D.
potentials revealed that DENV2 has a basic platform beneath its fusion loops, providing a possible means for its interaction with anionic lipids. Sphingolipids are also negatively charged, but it is less clear how they could interact with basic regions on SFV E1. Alpha- and flaviviruses also vary in pH requirements for efficient fusion, ranging from pH of ~6.2 (TBEV and SFV) (79, 80) to ~6.8 (DENV2) (81). It has been reported that flaviviruses with increased thermostability more efficiently infect certain hosts (82), and so it is conceivable that changes in optimal fusion pH might similarly affect transmission or tropism.

Comparative analysis of envelope structures from several viral serocomplexes and families provides valuable insights toward the understanding of class II fusion. The structure of SLEV E highlights several emerging serocomplex-specific structural features of E proteins. An understanding of the unique properties of these proteins should pave the way for structure-based experiments to illuminate the biochemical basis for flavivirus interactions with host determinants.

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