Reptilian Reovirus Utilizes a Small Type III Protein with an External Myristylated Amino Terminus To Mediate Cell-Cell Fusion

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Reptilian reovirus is one of a limited number of nonenveloped viruses that are capable of inducing cell-cell fusion. A small, hydrophobic, basic, 125-amino-acid fusion protein encoded by the first open reading frame of a bicistronic viral mRNA is responsible for this fusion activity. Sequence comparisons to previously characterized reovirus fusion proteins indicated that p14 represents a new member of the fusion-associated small transmembrane (FAST) protein family. Topological analysis revealed that p14 is a representative of a minor subset of integral membrane proteins, the type III proteins N_{exo}\!/C_{cyt} (N_{exo}/C_{cyt}), that lack a cleavable signal sequence and use an internal reverse signal-anchor sequence to direct membrane insertion and protein topology. This topology results in the unexpected, cotranslational translocation of the essential myristylated N-terminal domain of p14 across the cell membrane. The topology and structural motifs present in this novel reovirus membrane fusion protein further accentuate the diversity and unusual properties of the FAST protein family and clearly indicate that the FAST proteins represent a third distinct class of viral membrane fusion proteins.

Biological membrane fusion is an essential cellular process mediated by specific fusion proteins (22, 57, 61). Extensive analysis of a number of enveloped virus fusion proteins has contributed to a model of protein-mediated membrane fusion. Enveloped virus fusion proteins are complex, multimeric, type I N_{exo}\!/C_{cyt} integral membrane proteins that facilitate virus entry into cells by mediating fusion between the viral envelope and the target cell membrane. Two distinct classes of enveloped virus fusion proteins have been identified: the class I fusion proteins exemplified by influenza virus and human immunodeficiency virus proteins and the class II proteins of the alpha- and flaviviruses (25, 47, 56, 57). For both classes, triggered conformational changes and/or multimer reorganization of their complex ectodomains are essential aspects of the fusion reaction (25, 56). This transition from a metastable to a low-energy form is believed to provide the energy to overcome the thermodynamic barriers that inhibit spontaneous membrane mergers (30, 56). However, the necessity and/or precise role of structural remodeling as a thermodynamic mediator of the fusion reaction remains unresolved (3, 13, 14, 37).

Since nonenveloped viruses lack a lipid bilayer, virus entry is not dependent on membrane fusion. As a result, nonenveloped viruses do not encode membrane fusion proteins. The rare exceptions to this generalization are the fusogenic reoviruses, an unusual group of syncytium-inducing nonenveloped viruses, the reovirus fusion proteins are nonstructural nature and restricted role of the FAST proteins in the virus replication cycle may account for their small size and possible evolution toward the minimal protein determinants required to promote fusion of biological membranes.

The small size of the reovirus FAST proteins makes it difficult to envision how extensive conformational changes could play a role in either regulating the exposure of a buried fusion peptide or providing sufficient energy to overcome the thermodynamic barriers that maintain membrane structure.

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sequently, the reovirus FAST proteins are unlikely to adhere to the current paradigm of protein-mediated membrane fusion that has emerged from studies of the enveloped virus fusion proteins (22, 47, 61). It seems likely that an analysis of each individual member of the FAST protein family, followed by a comparison of the motifs required for their fusion function, will contribute to an improved mechanistic understanding of these rudimentary membrane fusion machines. We recently characterized a python reovirus, the prototype of a new species of fusogenic reovirus, reptilian reovirus (RRV) (1, 12). We now show that a p14 protein encoded by the first open reading frame (ORF) of a bicistronic mRNA represents a third distinct member of the reovirus FAST protein family with its own signature arrangement of structural motifs. Biochemical analysis revealed that p14 is a surface-localized, type III integral membrane protein (i.e., it utilizes an internal reverse signal-anchor sequence to direct an N_term/C_cyt membrane topology). This topology results in the cotranslational translocation of a small, myristylated ectodomain across the lipid bilayer. Although the precise role of the external myristylated N terminus of p14 in the membrane fusion reaction is undetermined, this discovery adds a new element to be considered in models of protein-mediated membrane fusion.

MATERIALS AND METHODS

Cells and virus. RRV was isolated from a python (Python regius) (1) and obtained from W. Ahne (University of Munich, Munich, Germany). Vero cells and quail QM5 cells (11) were maintained at 37°C in a 5% CO2 atmosphere in medium 199 with Earle’s salts containing 100 U of penicillin and streptomycin per ml and 5% or 10% heat-inactivated fetal bovine serum, respectively.

Plasmids, cloning, and sequencing. The procedure for cDNA synthesis and cloning of the RRV genome segments is described in detail elsewhere (9). The full-length S1 genome segment indicated that either or both of the S1 genome segment is functionally redundant, this discovery adds a new element to be considered in models of protein-mediated membrane fusion.

RESULTS

In vitro transcription and translation. The full-length S1 genome segment, p14 ORF, p14-2HAC, and p14 constructs containing site-specific substitutions were transcribed and translated in vitro by use of nucleoside-treated rabbit reticulocyte lysates (Promega). Translation products were characterized by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) using 15% acrylamide gels, and proteins were visualized by fluorography as previously described (7).

Fluorescent cell staining. Cells were seeded in culture plates containing coverslips and transfected 20 to 24 h after seeding as described above. At 6 h posttransfection, cell monolayers were washed twice with HBSS and either fixed with ice-cold methanol for staining of intracellular p14 or surface stained. Permeabilized cells were preblocked with whole goat or rat IgG (1:1,000) in HBSS for 30 min at room temperature, and then a primary antibody (1:800 rabbit polyclonal anti-p14 or 1:200 mouse monoclonal [IgG2b] anti-HA antibody) was adsorbed to cells for 45 to 60 min at room temperature. After primary antibody binding, cell monolayers were washed six times in HBSS, with a second centrifugation at 100,000 g for 1 h to recover integral membrane proteins. All soluble and membrane fractions were then immunoprecipitated and analyzed by SDS-PAGE as described above.

Fluorescence-activated cell sorting. Cell line monolayers were washed twice with HBSS and either fixed with ice-cold methanol for staining of intracellular p14 or surface stained. Permeabilized cells were preblocked with whole goat or rat IgG (1:1,000) in HBSS for 30 min at room temperature, and then a primary antibody (1:800 rabbit polyclonal anti-p14 or 1:200 mouse monoclonal [IgG2b] anti-HA in blocking buffer) was adsorbed to cells and washed as described above. After the addition of the substrate BCIP/NBT (5-bromo-4-chloro-3-indolyolphosphate/nitroblue tetrazolium) to allow color development, cells were visualized under a Nikon Diaphot inverted microscope at a magnification of ×200. Image-Pro Plus software (v. 4.0) was used to capture images of stained cells. The relative abundance of various p14 mutants to mediate syncytium formation was quantitated by a syncytial index assay. The numbers of syncytial foci and syncytial nuclei present in five random fields of view were determined by microscopic examination of Giemsa-stained transfected cell monolayers at ×100 magnification. Results were reported as the means ± standard errors from three separate experiments. For antibody inhibition studies, twofold serial dilutions of complement-activated polyclonal anti-p14 antisera or normal rabbit serum were added to the medium of p14-transfected cells at 4 h posttransfection. At 14 to 18 h posttransfection, the cell monolayers were fixed with methanol, stained with Wright-Giemsa stain, and examined for the presence of multinucleated syncytia.

The first ORF of the bicistronic S1 genome segment of RRV is responsible for cell-cell fusion. Sequence analysis identified two sequential, partially overlapping ORFs in the S1 genome segment of RRV (Fig. 1A). In vitro transcription and translation confirmed that the S1 genome segment is functionally bicistronic and encodes two protein products, namely a 35-kDa homolog of the ARV cell attachment protein eC2 and a 14-kDa RRV-specific gene product (Fig. 1B). Transfection analysis with the full-length S1 genome segment indicated that either or
both of these S1 gene products induce cell-cell fusion (Fig. 1C). Subcloning and expression revealed that the 125-amino-acid p14 protein encoded by the first ORF, when expressed by itself in transfected cells, induced extensive multinucleated syncytium formation in both transfected Vero epithelial cells and QM5 quail cell fibroblasts (Fig. 1C). Immunoprecipitation confirmed the expression of p14 in both transfected and virus-infected cells (Fig. 1B). Gapped alignments of p14 with the other FAST proteins of ARV, Nelson Bay reovirus, and BRV revealed no significant sequence similarity (percent amino acid identities of <15%) (data not shown), indicating that the RRV p14 protein is a new member of the FAST protein family.

**Structural motifs in p14.** The RRV p14 protein contains several predicted structural motifs (Fig. 2). A hydropathy plot of p14 and sequence analysis identified a predicted transmembrane (TM) domain, suggesting that p14 resides as an integral membrane protein. Sequence analysis also showed that p14 lacks a cleavable N-terminal signal sequence (29), suggesting that the membrane-spanning domain may function as an internal signal anchor (21). The only other region in p14 with any hydrophobic character occurs in the N-terminal domain (Fig. 2A), a region we termed the hydrophobic patch. The N-terminal domain also contains a consensus sequence for N-terminal myristylation (MGXXXT/S/T/A) (52). The C-terminal domain is comprised of two different regions, namely a highly basic, membrane-proximal region (10 of the 22 residues immediately following the TM domain are basic) and a C-terminal proline-
rich region (eight prolines between residues 99 and 112) that includes a stretch of five consecutive prolines. The C-terminal domain also contains a consensus sequence for N-linked glycosylation (NXS/T) at Asn121. The functional significance of these sequence-predicted structural motifs was further investigated.

**p14 assumes an N<sub>exo</sub>/C<sub>cyt</sub> surface membrane topology.** Analysis of the soluble and membrane fractions from transfected QM5 cells indicated that p14 localizes exclusively to the membrane pellet (Fig. 3A), suggesting that p14 is cotranslationally inserted into cellular membranes. Treatment of the membrane pellet with either high salt or high pH to extract peripheral membrane proteins (15) did not alter the p14 distribution (Fig. 3A), indicating that the predicted p14 transmembrane domain is functional and that p14 exists exclusively as an integral membrane-spanning protein. Consistent with the membrane localization of p14, immunostaining of permeabilized cells revealed a reticular staining pattern, with concentrations of p14 in the perinuclear region and numerous punctate foci throughout the cytoplasm radiating out to the plasma membrane. Similar staining of nonpermeabilized cells showed a patchy ring fluorescence at the surfaces of cells (Fig. 3B). Furthermore, a polyclonal anti-p14 antiserum inhibited syncytium formation (Fig. 3C). These results suggest that p14 localizes to the endoplasmic reticulum (ER)-Golgi pathway and that at least a portion of p14 traffics to the cell surface, where it is directly involved in promoting the membrane fusion reaction.

For examination of the p14 membrane topology, constructs were created that contained two HA epitope tags either added to the C terminus of p14 or inserted between residues seven and eight within the N-terminal domain. The addition of the double-epitope tag to the C terminus (p14-2HAC) slowed, but did not inhibit, the extent of cell-cell fusion, while insertion of the epitope tag in the N-terminal domain (p14-2HAN) abolished polykaryon formation (Fig. 4A). Cells transfected with p14-2HAN or p14-2HAC were immunostained with an anti-HA monoclonal antibody, either after fixation and permeabilization to reveal intracellular fluorescence or with live cells to detect the surface-expressed ectodomain. In permeabilized cells, both constructs revealed the characteristic reticular staining pattern of authentic p14 (Fig. 4B). Positive surface staining was only obtained with the p14-2HAN construct, indicating that p14 assumes an N<sub>exo</sub>/C<sub>cyt</sub> surface topology in the plasma membrane. Furthermore, the potential N-linked glycosylation site near the C terminus of p14 was nonfunctional (Fig. 5), as evidenced by the lack of a gel mobility shift due to glycosylation when p14 was expressed in transfected cells in the absence or presence of the glycosylation inhibitor tunicamycin. Therefore, we infer that p14 assumes an exclusive N<sub>exo</sub>/C<sub>cyt</sub> topology.

**The C terminus of p14 is dispensable for fusion activity.** Proline-rich regions can form a type II polyproline helix and are often involved in protein-protein interactions (36). For an examination of the influence of the unusual C-proximal polyproline motif on p14 fusion activity, five p14 C-terminal deletion mutants (named for their lengths, in amino acids) were assessed by a quantitative fusion assay based on the average number of syncytial nuclei per field. Deletion of the C-terminal 10 to 20 amino acids of p14, including the five consecutive proline residues and the potential polyproline helix (residues 108 to 112), reduced the rate of p14-induced syncytium formation by the anti-p14 antiserum. Scale bar = 100 μm.
syncytium formation, as evidenced by a decreased syncytial index at early times posttransfection (Fig. 6A). However, the overall extent of fusion remained unimpaired, as both p14-C105 and p14-C115 mediated the formation of large multinucleated syncytia (Fig. 6B) that eventually encompassed the entire cell monolayer. The p14-C88 deletion, which removed all of the proline residues from the C-terminal domain, also displayed an impaired rate, but not extent, of fusion (Fig. 6). Therefore, the C-terminal 37 amino acids of p14, representing the proline-rich region and potential polyproline helix motif, are dispensable for the mechanism of p14-mediated fusion.

Conversely, the p14-C78 construct was devoid of fusion activity, while the p14-C83 construct retained minimal fusion activity, initiating the formation of small syncytia that failed to progress in size over time (Fig. 6). The different fusion activities of the C-terminal truncations did not correlate with their relative expression-detection levels or efficiencies of membrane insertion; all of the truncated p14 constructs localized exclusively to the membrane fraction (Fig. 6C). Immunoprecipitation results did indicate a reduced detection of the C88, C83, and C78 constructs in the membrane fraction, possibly due to decreased protein expression or stability and/or the loss of a C-proximal epitope recognized by the polyclonal antiserum (Fig. 6C). However, the relative fusion activity of these constructs did not correlate with their detection in the membrane fraction. Therefore, a minimum size or structure of the p14 C-terminal domain most likely influences p14 fusion activity by altering events downstream of membrane insertion.

N-terminal myristylation of p14 is essential for fusion activity. The myristate moiety of myristylated proteins is almost always associated with the cytoplasmic leaflet of lipid bilayers (38). In view of the predicted N_exo/C_cyt topology of p14, we anticipated that the myristylation consensus sequence would be nonfunctional and therefore irrelevant for the p14 function. This was not the case. Radiolabeling indicated the incorporation of [3H]myristic acid into authentic p14, but not into a construct containing a G2A substitution that removed the myristylation consensus signal (Fig. 7A). The loss of labeling of the G2A construct by myristic acid confirmed that the labeling of authentic p14 reflected the incorporation of [3H]myristic acid and not the metabolic redistribution of the radiolabel, and it implied that p14 is a myristylated integral
membrane protein. In addition, the myristate moiety appears to be an essential component of p14, since the p14-G2A construct did not mediate cell-cell fusion (Fig. 7B). The altered fusion activity of the myristylation-negative construct did not reflect altered protein expression (Fig. 7A) or p14 subcellular localization, as determined by immunofluorescent staining (Fig. 7C). Therefore, the N-terminal myristylation consensus sequence of p14 is both functional and essential for p14-induced syncytium formation.

p14 translocates its myristylated N terminus across the membrane. The unexpected placement of a myristylated N-terminal domain external to the plasma membrane prompted the need for further confirmation of the p14 topology. An N-linked glycosylation site was engineered into the N-terminal domain of p14 (V9T substitution), and N-glycosylation mapping was used to examine the topology of the p14-V9T construct. In vitro translation of authentic p14 in the presence or absence of canine microsomal membranes (CMM) produced polypeptides with identical gel mobilities (Fig. 8A), confirming the sequence-predicted absence of a cleavable N-terminal signal peptide. Conversely, in vitro translation of p14-V9T in the presence of CMM resulted in two protein species, one with a retarded gel mobility relative to authentic p14 or to p14-V9T translated in the absence of CMM (Fig. 8A). The mobility shift of approximately 3 kDa approximated the expected shift per core glycan (2.5 kDa) (6) and was in accord with the mobility shift of a known glycosylated control protein (Fig. 8A). The nonmyristylated p14-G2A construct in the presence of CMM demonstrated a minor population with the same gel mobility as the glycosylated species of p14-V9T (Fig. 8A), suggesting that a small percentage of p14-G2A molecules may adopt the inverse Ncyt/Cexo topology, leading to the glycosylation of Asn121. Therefore, myristylation may be a minor contributor for determining an exclusive Nexo/Ccyt topology for p14, at least in vitro.

Analysis of p14-V9T in transfected Vero cells also detected two different species of p14-V9T, with a loss of the slower migrating form after the treatment of cells with tunicamycin, confirming that this polypeptide represented glycosylated p14 (Fig. 8B). To exclude the possibility that there were two populations of p14, one with the myristylated N-terminal domain inside the cell and the other nonmyristylated form in the Nexo/Ccyt topology, p14-V9T-transfected cells were labeled with [3H]myristic acid. The N-terminal domain of p14-V9T was both N-glycosylated and myristylated, as evidenced by the ability of [3H]myristic acid to label both forms of p14-V9T (Fig. 8B). These results conclusively establish that the myristylated N-terminal domain of p14 is translocated into the ER lumen, where we believe it interacts with the luminal leaflet of the membrane, and after trafficking to the plasma membrane, resides in the exoplasmic leaflet outside the cell.

An additional observation noted during in vivo analysis was...
the loss of syncytium-inducing ability imparted by the V9T substitution (data not shown). The loss of fusion activity was not the result of steric hindrance from the addition of a large carbohydrate moiety, since the fusion ability of p14-V9T was not restored by the inhibition of N-linked glycosylation by tunicamycin. Transport of the p14-V9T construct was also not affected, as a population of p14-V9T molecules was clearly visualized accumulating at the plasma membrane by immunofluorescence staining of permeabilized cells (data not shown).

Therefore, a minor alteration near the N terminus of the p14 ectodomain hydrophobic patch affects the membrane fusion activity of the protein, independent of any effects on p14 membrane insertion, protein topology, or cell surface localization. The small p10 ectodomain contains a similar hydrophobic patch that may function in an analogous manner as the fusion peptide motifs found in enveloped virus fusion proteins (45). However, the relatively low overall hydrophobicity of these FAST protein hydrophobic patches and the likelihood that these potential fusion peptide motifs are not sequestered within a complex tertiary structure distinguish these motifs from the typical fusion peptides of enveloped viruses (47, 57).

**FIG. 7.** N-terminal myristylation of p14 is essential for cell fusion. (A) 2HAC-tagged p14, p14-G2A, and pcDNA3-transfected Vero cells were labeled with either [3H]myristic acid (Myr.) or [3H]leucine (Leu.) and immunoprecipitated with anti-p14 polyclonal antiserum (α p14) or normal rabbit serum (nrs). Precipitates were fractionated by SDS-PAGE and radiolabeled p14 was detected by fluorography. (B) Vero cells transfected with either 2HAC-tagged p14 (a) or p14-G2A (b) were fixed with methanol at 18 h posttransfection and immunostained with anti-p14 polyclonal antiserum and an alkaline phosphatase-conjugated secondary antibody. Arrows indicate the limits of an antigen-positive syncytial focus (a) or individual antigen-positive cells expressing the fusion-negative p14-G2A construct (b). Scale bar = 100 μm. (C) QM5 cells transfected with p14-G2A were permeabilized with methanol and immunostained at 9 h posttransfection with anti-p14 polyclonal antiserum and FITC-conjugated secondary antibody. Fluorescence microscopy (a) revealed punctate, perinuclear intracellular staining of p14-G2A and a ring surface fluorescence characteristic of authentic p14. The corresponding DIC image overlaid with the fluorescent image is also shown (b). Scale bar = 10 μm.

**FIG. 8.** p14 translocates its myristylated N terminus across the membrane. (A) 2HAC-tagged p14, p14-G2A, or p14-V9T were in vitro transcribed and translated in the presence or absence of canine microsomal membranes (CMM). The relative gel mobilities of the radiolabeled authentic (p14) and glycosylated (p14*) species of p14 are indicated on the right, and molecular weight markers are shown on the left. The control lane (Con.) indicates the glycosylated and nonglycosylated species generated by the translation of yeast α factor mRNA. (B) Vero cells were transfected with 2HAC-tagged p14 or p14-V9T or were mock transfected (M) with pcDNA3 vector. Transfected cells were labeled with [3H]myristic acid (Myr.) or [3H]leucine (Leu.) in the presence or absence of tunicamycin (tunic.) and were immunoprecipitated with anti-p14 antiserum (α p14) or normal rabbit serum (nrs). The relative gel mobilities of the radiolabeled authentic (p14) and glycosylated (p14*) species of p14 are indicated on the right, and molecular weight markers are shown on the left.

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**DISCUSSION**

Structural motifs in p14, the newest member of the diverse FAST protein family. The present study indicates that the RRV p14 protein is a novel reovirus FAST protein and the newest member of a diverse family of atypical viral membrane fusion proteins. While p14 shares no significant sequence sim-
proteins include cytochrome P-450, CLN3, mouse synaptogamin I and II, and the influenza viral proteins M2 and NB (24, 34, 35). Other examples of type III membrane proteins are also myristylated at internal lysine or cysteine residues (2, 18, 49, 50, 53). In the case of myristylated integral membrane proteins, stable membrane association derives from the TM domain and the myristic acid serves some undefined function other than protein anchoring in the membrane. The role of myristate in these integral membrane proteins is speculative, but it has been suggested to involve the regulation of enzymatic activity by the modulation of protein-protein or protein-lipid interactions or by an alteration of the ability of the protein to be subsequently modified.

Our discovery that the myristylated p14 N terminus is translocated across the membrane and resides outside the cell was surprising, but not without precedent. The large surface antigens of both human and duck hepatitis B viruses are polytopic proteins that adopt two functional membrane topologies, one of which externalizes the myristylated N-terminal pre-S1 domain with cytosolic Hsc70 lead to posttranslational or an electrostatic interaction (33). The unique ability of myristic acid to reversibly associate with the cytosolic leaflet of the plasma membrane in response to the availability of a second signal regulates the activities of a number of cellular signaling proteins (35, 38, 41).

In addition to soluble proteins, a limited number of integral membrane proteins are N-terminally myristylated, such as the type II (Cexo/Ncyt) CLA-1 protein (CD36 family) (5). Some integral membrane proteins are also myristylated at internal lysine or cysteine residues (2, 18, 49, 50, 53). In the case of myristylated integral membrane proteins, stable membrane association derives from the TM domain and the myristic acid serves some undefined function other than protein anchoring in the membrane. The role of myristate in these integral membrane proteins is speculative, but it has been suggested to involve the regulation of enzymatic activity by the modulation of protein-protein or protein-lipid interactions or by an alteration of the ability of the protein to be subsequently modified.

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are functional and whether additional examples of myristylated ectodomains exist.

A myristylated ectodomain is essential for p14 fusion activity. The type II topology (C_{exo}/N_{cyt}) of N-terminally myristylated cellular membrane proteins or the locations of the internally myristylated lysine or cysteine residues in these proteins position the myristic acid on the cytosolic side of the membrane, similar to the situation with myristylated cytosolic proteins. In contrast, the essential myristic moiety present in p14 must exert its influence on p14-induced membrane fusion via interactions with the external leafllet of the membrane bilayer. Although the role of the p14 myristic acid is unclear, several possible roles can be envisioned. Acylation of both viral and cellular proteins can target these proteins to detergent-resistant membrane microdomains within the plasma membrane (31, 39). Although the myristylation-negative p14-G2A still trafficked to the cell surface (Fig. 5), the loss of myristylation could influence p14 localization to membrane microdomains. A second possibility is based on the assumption that the myristic acid associates with the luminal leafllet of the ER membrane after translocation, in a manner analogous to the membrane association of signal peptides before their cleavage (19, 20). Signal peptide interactions with the luminal leafllet of the ER membrane can promote the correct folding of the nascent protein (59); a similar situation could contribute to folding of the p14 ectodomain into a fusion-competent conformation. An intriguing third possibility reflects the unique ability of myristic acid to reversibly associate with a lipid bilayer (38). During the close apposition of membranes that must occur prior to fusion, myristic acid may disassociate from the outer leafllet of the donor membrane and interact with the outer leafllet of the target membrane. Such interactions of myristic acid, from a sufficient number of p14 molecules, with the target and/or donor membranes could alter the lipid packing of one or both membranes to support the lipid rearrangements required for fusion to proceed. This hypothesis is supported by the greater membrane permeabilization exhibited by a myristylated hydrophobic peptide than by its nonacylated partner (23).

Our discovery of this newest FAST protein with its unusual membrane topology further underscores the diversity of class III viral membrane fusion proteins. A collection of membrane interaction motifs, including signal-anchors, hydrophobic patches, polybasic regions, and now an externalized fatty acid, have been assembled into these rudimentary membrane fusion proteins. Though the mechanism of FAST-mediated membrane fusion remains undetermined, it is evident that the FAST proteins do not have the size capability or structural features to promote membrane fusion by using large energy-releasing triggered conformational changes, as occurs with the class I and class II enveloped virus fusion proteins (47). Continued structure-function analysis of the FAST proteins should provide alternate insights into the minimal determinants of protein-mediated fusion of biological membranes.

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