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To enter cells, enveloped viruses use fusion-mediating glycoproteins to facilitate the merger of the viral and host cell membranes. These glycoproteins undergo large-scale irreversible refolding during membrane fusion. The paramyxovirus parainfluenza virus 5 mediates membrane merger through its fusion protein (F). The transmembrane (TM) domains of viral fusion proteins are typically required for fusion. The TM domain of F is particularly interesting in that it is potentially unusually long; multiple calculations suggest a TM helix length between 25 and 48 residues. Oxidative cross-linking of single-cysteine substitutions indicates the F TM trimer forms a helical bundle within the membrane. To assess the functional role of the paramyxovirus parainfluenza virus 5 F protein TM domain, alanine scanning mutagenesis was performed. Two residues located in the outer leaflet of the bilayer are critical for fusion. Multiple amino acid substitutions at these positions indicate the physical properties of the side chain play a critical role in supporting or blocking fusion. Analysis of intermediate steps in F protein refolding indicated that the mutants were not trapped at the open stalk intermediate or the prehairpin intermediate. Incorporation of a known F protein destabilizing mutation that causes a hyperfusogenic phenotype restored fusion activity to the mutants. Further, altering the curvature of the lipid bilayer by addition of oleic acid promoted fusion of the F protein mutants. In aggregate, these data indicate that the TM domain plays a functional role in fusion beyond merely anchoring the protein in the viral envelope and that it can affect the structures and steady-state concentrations of the various conformational intermediates en route to the final postfusion state. We suggest that the unusual length of this TM helix might allow it to serve as a template for formation of or specifically stabilize the lipid stalk intermediate in fusion.
Introduction

Membrane fusion is a fundamental biological process that occurs in intracellular trafficking, exocytosis, rescaling of plasma membranes, protein trafficking, and in the entry of enveloped viruses. This ubiquitous process is mediated and tightly controlled by a combination of specific protein machinery and lipid composition. Lipids spontaneously assemble into bilayer structures such as liposomes; however, lipid bilayer membranes do not spontaneously fuse. The spontaneous negative or positive curvature of a lipid can enhance or diminish fusion, respectively, but energy must be expended to overcome hydration repulsion between membranes and to disrupt the bilayer structure. The energy for this remodeling may be derived from the thermal fluctuations of the membrane or from specialized fusion proteins. Many biological processes, such as neuronal synaptic vesicle fusion, endosomal fusion, and exocytosis, employ the SNARE superfamily proteins. SNAREs are found in all eukaryotic organisms. All SNARE proteins have a common heptad repeat that forms four-helix coiled-coil structures, and this coiled-coil SNARE complex forms in trans to promote fusion of the two membranes in which the SNARE proteins are anchored. Enveloped viruses use an analogous strategy and mediate fusion with target cells through specialized fusion proteins. The paramyxovirus parainfluenza virus 5 (PIV5) requires two surface glycoproteins for this process: the attachment protein hemagglutinin neuraminidase (HN) that binds sialic acid and the fusion protein (F) that physically merges the two membranes. Paramyxovirus fusion occurs at the plasma membrane and does not require the low pH of the endosome to trigger fusion.

The paramyxovirus F protein is a class I fusion glycoprotein that is synthesized as a type I integral membrane protein and it folds into homotrimers, is posttranslationally modified by the addition of carbohydrate chains, and then is proteolytically cleaved to become biologically active. Similar processing occurs for other class I viral fusion proteins, such as influenza virus hemagglutinin (HA), HIV gp160, retrovirus Env, Ebola GP, and SARS CoV S. The paramyxovirus F precursor protein (F0) is cleaved into the membrane-anchored F1 subunit and the smaller N-terminal F2 fragment. F1 contains two hydrophobic regions, the N-terminal fusion peptide (FP), located at the new N-terminus after cleavage, and the transmembrane (TM) domain, and two heptad repeat regions, HRA and HRB. HRA is located immediately C-terminal to the FP, and HRB is proximal to the TM domain.

The paramyxovirus F protein folds initially into a metastable prefusion form (Fig. 1a) that upon triggering undergoes a series of large scale conformational rearrangements, proceeding down an energy gradient to form a final irreversible postfusion form (Fig. 1f).
Recently, the crystal structures of both the uncleaved prefusion conformation of the paramyxovirus F protein and the uncleaved postfusion conformation were solved.\textsuperscript{6,7} The prefusion form contains a globular head containing three domains (DI–DIII) attached to a trimeric coiled-coil stalk formed by the HRB region. The HRA region in the prefusion form is composed of 11 distinct segments that wrap around the DIII core in the globular head.\textsuperscript{7} This is in contrast to the postfusion form of F where HRA is extended into a long α-helix as part of the six-helix bundle (6-HB). For the postfusion structure, an unanticipated finding emerged as the available data indicate the F TM domain and/or cytoplasmic tail are important for the folding of F into the metastable prefusion form of F.\textsuperscript{6} Secreted F lacking a TM domain converts to the postfusion form.

Upon receptor binding, biochemical studies indicate HN induces a conformational change in F and the HRB three-helix stalk separates.\textsuperscript{8} It is hypothesized that following the melting of the HRB helices and destabilization of the head (Fig. 1b), HRA refolds to form an extended α-helical coiled coil, which enables the insertion of the fusion peptide into the target cell membrane and forms the prehairpin intermediate (Fig. 1c). The F protein then refolds where HRB binds into the grooves between the HRA regions and forms the six-helix bundle (6-HB), bringing the FP and TM domain into juxtaposition within the same membrane\textsuperscript{7–9} (Fig. 1e). A conceptually related final postfusion structure is formed for all enveloped virus fusion proteins. The postfusion structure consists of either an α-helical coiled-coil structure like the 6-HB in PIV5 F or β-strand structures in other fusion proteins, such as dengue virus E, Semliki Forest virus E1, and vesicular stomatitis virus (VSV) G.\textsuperscript{12} In all cases the fusion peptide and TM domain are together in the same membrane. The formation of the 6-HB and the associated free-energy change is tightly linked to the merger of the target and viral membranes.\textsuperscript{9,13} The collapse of the prehairpin intermediate distorts the bilayers to possibly form an initial pointlike protrusion.\textsuperscript{14} Whether the insertion of the fusion peptide perturbs the lipid bilayer and lowers the distortion energy is unknown.\textsuperscript{14} Following the initial bilayer contact, membrane merger proceeds to the lipid stalk intermediate, where the outer leaflets of each bilayer merges but the inner leaflets remain separate. This stalk then expands to form the hemifusion diaphragm. Continuation of this expansion leads to formation of the fusion pore that allows for the transfer of aqueous contents (Fig. 1f–i).\textsuperscript{15}

In addition to aiding in protein folding and stability, the TM domains of many viral envelope fusion proteins have been shown to have a role in fusion.\textsuperscript{16–24} When the TM domain of influenza virus HA was replaced with a glycosylphosphatidylinositol anchor, it resulted in a fusion protein that could mediate hemifusion by allowing the transfer of a lipid but not aqueous fluorescent probe, suggesting a role of the TM domain in pore formation and enlargement. This glycosylphosphatidylinositol-anchored HA was embedded only in the outer leaflet of the membrane, and it was proposed that the TM domain, which spans the entire membrane, affects the positive curvature on the inner leaflet that would allow for pore formation.\textsuperscript{17,19} In addition, specific residues in the TM domain have been shown to be important for fusion of influenza virus HA,\textsuperscript{18} VSV G,\textsuperscript{20} HIV gp41,\textsuperscript{23–25} and baculovirus GP64.\textsuperscript{21}

Although little is known about the role of the TM domain in paramyxoviruses, the TM domain of PIV5 F shows several features that might be relevant to function.

\begin{align*}
\text{DIYLSAATSI TTSSVSLIG ICQSLGIL ILISVVWKL LLIUVRAI} & \\
\text{VYLYVSLGIL ICQSLGIL ILISVVWKL LLIUVRAI} & \\
\text{DYLSAATSI TTSSVSLG ICQSLGIL ILISVVWKL LLIUVRAI} & \\
\text{DYLSAATSI TTSSVSLG ICQSLGIL ILISVVWKL LLIUVRAI} & \\
\text{DYLSAATSI TTSSVSLG ICQSLGIL ILISVVWKL LLIUVRAI} & \\
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\text{DYLSAATSI TTSSVSLG ICQSLGIL ILISVVWKL LLIUVRAI} & \\
\text{DYLSAATSI TTSSVSLG ICQSLGIL ILISVVWKL LLIUVRAI} & \\
\end{align*}

It has a 25-residue hydrophobic segment (spanning from 485 to 509) that is somewhat longer than the typical hydrophobic length of 20 residues for a vertically inserted TM helix. The N-terminus of the predicted TM helix is devoid of side chains such as Trp, Tyr, Arg, or Lys that typically anchor the end of the helix in the head-group region of the bilayer, and the side chains of the 15 preceding residues are uncharged and can partition easily between bilayer and aqueous phases.\textsuperscript{26,27} This region forms the end of the coiled-coil HRB stalk and a short linker segment in the prefusion form of F. Finally, the C-terminal end of the TM helix is predicted to occur at the positively charged K510, but the seven residues following this residue are predominantly hydrophobic amino acids. Thus, if K510 snorkels into the head-group region, the TM helix might extend beyond this point. This finding is reminiscent of the TM helix of HIV gp41,\textsuperscript{23} which contains an absolutely conserved and functionally essential positively charged residue within the hydrophobic region of the TM helix. In summary, the TM helix of F appears to be longer than usual and has an unusually large number of potential orientations in membranes; these features piqued our interest and in part motivated the present study.

We first asked whether the TM domain formed a well-defined helical bundle in the prefusion form of the protein. This is a particularly important question, because TM helices of fusogenic proteins often have sequences known to mediate helix–helix association, such as the GXXXG motif, and mutation of these sequences sometimes, but not always, leads to proteins that are properly expressed but have impaired function.\textsuperscript{20,28,29} Cryo-electron microscopy (cryoEM) studies on the human immunodeficiency virus fusion protein (Env) differ, as some reports indicate the TM domains of the fusion protein are apart, like the legs of a tripod,\textsuperscript{30,31} whereas other studies indicate the TM domains are together and form one helical bundle.\textsuperscript{32} By singly substituting each residue of the TM domain with cysteines, we have examined the structure of the TM domain in the lipid membrane. The addition of an oxidative cross-linker and formation of disulfide bonds clearly showed that one face of the TM helix mediates the formation of a homomeric TM helical bundle, most likely a trimer. To address the role of specific residues within the TM
helix we performed alanine-scanning mutagenesis, and determined that two residues, 486 and 488, are critical for PIV5 F fusion. The aggregate of analysis of the steps of fusion indicate these mutants do not support hemifusion and are trapped at the lipid stalk stage just prior to formation of the hemifusion diaphragm. Our results indicate a specific amino acid sequence of the TM domain is necessary for completion of fusion. The amino acid side chains at residues 486 and 488, which are predicted to be in the outer leaflet of the F TM domain, are critical for the merger of the two lipid membranes and fusion completion. The block in fusion observed with mutation of F residues 486 and 488 could be overcome by addition of compounds that affect the curvature of the membrane bilayer.

Results

The F protein TM domains are in close proximity and likely form a three-helix bundle

The TM domain is not part of the solved crystal structure of either the prefusion form of PIV5 F° or the postfusion form of hPIV3 F. Although it is thought likely that the TM domain of F would be α-helical because the hydrophobic packing of an α-helix would aid in spanning a lipid bilayer, this has not been determined. It is also unknown if the TM domains of the PIV5 F monomers interact to form a trimer. It has been shown recently that TM peptides of influenza virus HA associate with each other in model membranes. In the prefusion form of F, HRB forms a three-helix bundle (3-HB) and HRB is separated from the TM domain by only seven residues. Therefore, it seems likely that the F TM domain monomers would be in close proximity in the membrane if not in a 3-HB. We used oxidative disulfide cross-linking to examine the structure of the F TM domain. This method has been used to investigate the arrangement of TM domains for several membrane proteins, such as the Escherichia coli chemoreceptor, the aspartate receptor, the influenza virus M2 ion channel, and CD39.

The PIV5 F protein contains 10 disulfide-bonded cysteine residues in the ectodomain and one free cysteine residue in the TM domain. This latter cysteine at residue 492 was mutated to serine, and single-cysteine substitutions were made in the TM domain in this pseudo wild-type (wt) Cys− background. All mutants in the Cys− background were expressed at the cell surface equivalently to pseudo wt F (data not shown). The fusion activity of these mutants was determined by using the luciferase reporter gene assay (Fig. 2). Most mutants showed fusion activity similar to that of pseudo wt (Cys−, C492S) and wt F protein (denoted as 492). Mutants 485, 486, 488, 497, and 503 exhibited some decrease in fusion activity, with the maximum reduction being ∼50% for mutant 486 (see Fig. 2). Mutations in the TM domain of influenza virus HA have been shown to affect raft association. However, none of the cysteine substitutions affected raft association of the F protein (data not shown).

We examined the effect of oxidative cross-linking of these F TM domain single-cysteine substitutions. For each mutant, there are three available cysteines in each trimer. If these cysteines are oriented toward each other in the membrane and are within disulfide bond-forming distance, two will form a disulfide bond and leave the third cysteine unbonded. On the other hand, if the helices are not specifically interacting, one might expect that high concentrations of oxidants would be required to induce any disulfide formation in the membrane and that there would be no clear pattern for forming disulfides between residues along any one face of the helix. In Fig. 3a are shown untreated F mutants analyzed under nonreducing conditions on a 3.5% SDS-PAGE gel. Disulfide bonds formed by air oxidation should occur only in the extracellular aqueous phase and the headgroup region of the outer leaflet of the bilayer. Air oxidation is much slower in the hydrophobic region of the bilayer, and the reducing intracellular environment minimizes air oxidation on the cytoplasmic side of the membrane. Cysteine residues in the outer leaflet
of the bilayer at positions 485, 486, 487, and 489 did form disulfide bonds, indicating the individual F TM domains are in close proximity to themselves and that these positions occur near the head-group region of the outer leaflet of the bilayer.

The data are in very good agreement with the expected orientations of the TM helices, as inferred from deletion mutagenesis and crystallographic analysis of the protein. In the prefusion structure of F residues 446 to 476 form a three-stranded coiled coil. Not in the atomic structure is a short linker that connects to the TM domain near residues 485. Four residues (481–484) can be deleted with full retention of function, while a two-residue deletion (483–484) caused a decrease in fusogenic activity. More strikingly, deletion of eight residues (477–484), which compose the end of the coiled coil and the entire linker, leads to nearly full retention of fusogenic
activity, while deletion of only six residues (479–484) leads to a protein that is less well-expressed on the cell surface and highly inactive in fusion.10 Interestingly, if the HRB coiled coil continues into the TM region, the four-residue and eight-residue mutants would position the side chains of residues 485 and 486 in close proximity, consistent with their high level of cross-linking in Fig. 3a (see also Supplemental Fig. 2). On the other hand, there would be a strong mismatch between the axial rotation of the helical axes in the HRB coiled coil versus the TM bundle in the six-residue deletion mutant, explaining its lack of stability and function.

Cu[(II)(1,10-phenanthroline)₃ (CuP) was used to induce disulfide cross-linking within the lipid interior,39,43 which is otherwise resistant to oxidation. When the F TM mutants were treated with 3 mM (final concentration) of CuP for 10 min at 37 °C, more extensive disulfide cross-linking was observed (Fig. 3b). Under conditions required to induce oxidation of the membrane-spanning regions, much more extensive cross-linking is observed in the aqueous-accessible region (residues 485–491). Within the TM region, residues 493, 494, 496, 498, 501, 504, 506, and 508 all showed disulfide bond formation. CuP (3 mM) was also added at 4, 10, and 22 °C for 10 min (data not shown). All residues near the ectodomain (485–491, 493, 494) formed disulfide bonds at all temperatures, even at 4 °C. Mutants that formed disulfide bonds after treatment with 3 mM CuP for 10 min at 37 °C also formed disulfide bonds when treatment with 3 mM CuP was for 1 or 5 min (data not shown). The effect of cross-linking on fusion could not be determined after oxidation due to the toxicity of the CuP on the live cells that are required for the fusion assays (data not shown).

Cross-linking occurs in a pattern suggestive of that of a helical bundle. Even at a qualitative level, it is clear that the extent of cross-linking follows the same periodicity as that of an α-helix (approximately 3.6 residues per turn), suggesting that one face of the helix interacts in the core of a bundle. Furthermore, the extent of cross-linking under these conditions is similar to that observed with previously characterized membrane proteins of known three-dimensional structure, suggesting that the TM domain folds into a well-defined structure.38,43 One unusual feature, however, is that three consecutive residues near the outer leaflet of the bilayer (489–491) form cross-links quite efficiently, suggesting some malleability in this region of the protein. To better determine the nature of the interface, the extent of disulfide cross-linking was quantified and fit to a sine wave. The raw CuP cross-linking data were normalized by dividing the amount of disulfide formed by the total amount of protein (disulfide linked plus not linked). The CuP data at 37 °C were used for most of the TM domain, positions 492–509. Because cross-linking was stronger in the outer leaflet of the bilayer, low-temperature (4 °C) CuP data were used for residue positions 485–491 so that data from the two regions would be on the same scale (and to maintain linearity of the image plate response to radioactivity). This type of approach was previously used to model the structure of the TM domain of the influenza virus M2 protein,38 where a similar level of cross-linking was observed.

It was considered that the disulfide cross-linking could occur between two trimers. However, when more than 100-fold different amounts of F protein were expressed, there was no change in the level of cross-linking as would be expected to occur for intertrimer cross-links (data not shown). In addition, higher molecular weight species of trimer–trimer formation, in addition to cross-links within a trimer, were not observed on gels (data not shown). Further, the extensive nature of the cross-links for some mutants and not others argues for cross-links within the TM domains within a trimer.

Examination of the F TM domain data showed that the cross-linking efficiency oscillated as a function of the sequence position with approximately the same periodicity as the α-helix (3.6 residues per turn). However, G497 deviated from this trend, showing a lower degree of cross-linking than expected. Given the unusual characteristics of glycine residues, residue 497 was removed from the calculation and separate fits were made to the data before and after this residue. The periodicity was also indicative of a helical conformation (residues 485–496: 3.65±0.18 residues per turn; residues 498–509: 3.40±0.21 residues per turn).

Fig. 3. F protein TM domain cysteine substitution and disulfide bond formation on oxidation. (a) F protein TM domain cysteine mutants in a pseudo wt background (Cys−, C492S) were expressed in HeLa CD4 LTR βgal cells, and 18 h.p.t. cells were transfected and labeled as above. Dounce homogenized, and treated with 3 mM CuP for 10 min at 37 °C. Samples were then solubilized, immunoprecipitated with an anti-F2 PAb. Polypeptides were analyzed by SDS-PAGE on a 3.5% acrylamide gel under nonreducing conditions. (b) Cells were transfected and labeled as above. Dounce homogenized, and treated with 3 mM CuP for 10 min at 37 °C. Samples were then solubilized, immunoprecipitated with an anti-F2 PAb, and polypeptides were analyzed by SDS-PAGE on a 3.5% acrylamide gel under nonreducing conditions. Circled residues indicate interacting residues in the predicted helix. (c) Periodicity of the predicted TM helix. The raw CuP cross-linking data in (b) were normalized by dividing the amount of disulfide formed by the total amount of protein. Also shown is the sine wave fit of the data. The data obtained with oxidation occurring at 4 °C were used for residue positions 485–491, and the data obtained with oxidation occurring at 37 °C were used for the remaining positions (492–509). This was done to compensate for the greater disulfide bond formation that takes place in the outer leaflet and to maintain the signal on the image plate used to detect radioactivity within the linear range. (d) Ideal helix (left) and pi bulge (right) models of the TM helices. Interface positions predicted from disulfide cross-linking are shown in green spheres. For the first model, the TM domain is predicted to form a helix with a pi bulge at residues 497–500 (arrow). (e) Model of the TM helix trimer containing a pi bulge (in green). On the left is a view from within the membrane and on the right is a view from above. As in (d), interface positions are shown in green spheres. (f) Model of the TM trimer containing a stutter (in red). On the left is a view from within the membrane and on the right is a view from above. Interface C° positions for the model are shown in red spheres (C° shown for G497).
residues per turn; Fig. 3c). The pattern is interrupted by the insertion of one residue near residue 497. Thus, the fit parameters suggest a packed helical TM interface made up of residues 486, 489, 493, 501, 504, and 508 (circled in Fig. 3b) with a potential distortion to the helix around residue 497.

To create models for the TM region, we assume it forms a trimeric bundle and that efficient cross-linking occurred only at residues close together in the three-dimensional structure. One way to allow insertion of a single residue into α-helices is by inserting a pi bulge.44,45 Figure 3d shows an example of a 3-fold symmetric bundle created by using the backbone of a TM helix containing a pi bulge from the Na+-dependent aspartate transporter structure (Protein Data Bank code 2nwl, 2.96 Å).46 The sequence of the TM region of F was threaded onto this backbone, and Gly497 was positioned within the pi bulge. The effect of a pi bulge on the positioning of the residues that cross-link efficiently (when mutated to Cys) can be seen by comparing it to the result expected for an ideal helix and without a pi bulge (Fig. 3d). The pi bulge allows both the interfaces above and below the bulge to occur on the same face of the helix. A bundle was created from this helix by applying C3 symmetry followed by energy minimization (Fig. 3e).

A second way insertions are accommodated in coiled coils is by variation of the helix packing in a common structural motif known as a “stutter.”47 A geometric model incorporating this insertion was created by correlating the cross-linking data with the Cα distances of the trimeric coiled coil from influenza virus HA. As the stutters in HA result in different local distortions,48 pre- and postfusion HA structures49,50 were searched for one that matched the cross-linking data as described in Supplemental Methods. Threading the sequence of the TM helix onto this backbone template gave a model in good qualitative agreement with the cross-linking data (Fig. 3f). In summary, the cross-linking data show qualitative agreement with the cross-linking data and 6-CF, were transferred en bloc to the CV-1 cells. Shown is the quantification of cell surface expression of the F protein mutants TM01–TM09 was equivalent to wt PIV5 F (Table 1). However, mutant TM10 was not expressed at the cell surface; thus, F TM10 is likely to be a malfolded protein that is not transported through the exocytic pathway to the cell surface.51 Previously, when the TM domain of PIV5 HN was replaced with leucine residues, there was no discernable effect on the cell surface expression of HN and no effect on its biological function.52

The structure of the TM domain was taken as beginning at V485 and ending at W509. The alanine substitutions in the alanine scan were labeled with the lipidic probe R18 and the aqueous probe 6-CF. Labeled RBCs were bound to CV-1 cells for 1 h at 4 °C and then incubated at 37 °C for 15 min before visualization by confocal microscopy. Cell–cell fusion was observed as the transfer of red R18 and green 6-CF from the target RBCs to the effector CV-1 cells. Shown is the quantification of 6-CF. The means and error bars are from three microscopic fields.

**F TM domain residues L486 and I488 are key residues involved in fusion**

To determine further which residues in mutants TM01 and TM02 are responsible for the greatly reduced fusion activity, the first five residues, 485–489, of the F protein TM domain were changed individually to alanine. All these mutants were expressed at the cell surface at levels similar to that of wt F (Table 1). F TM domain mutants V485A, S487A, and I489A formed syncytia at levels similar to that of wt F, but F TM domain mutants L486A and I488A did not cause syncytia formation (Fig. 5a). These mutants also showed a substantial decrease in fusion in the luciferase reporter and dye transfer assays, TM01 and TM02 caused a consistent and major reduction in fusion in all assays used (Fig. 4c and d).
using F-specific mAb F1a. Surface expression levels were determined by flow cytometry using F-specific mAb F1a. The earliest stage of fusion

Table 1. Relative surface expression of mutant F proteins

| Mutant     | RMFI (% wt) |
|------------|-------------|
| wt         | 100         |
| TM01       | 102         |
| TM02       | 99          |
| TM03       | 103         |
| TM04       | 98          |
| TM05       | 94          |
| TM06       | 87          |
| TM07       | 90          |
| TM08       | 111         |
| TM09       | 97          |
| TM10       | 9           |
| V485A      | 102         |
| L486A      | 81          |
| S487A      | 99          |
| I488A      | 89          |
| I489A      | 99          |
| S443P      | 113         |
| S443P/L486A| 67          |
| G105A      | 97          |
| G105A/I488A| 78          |
| G105A/1488A| 85          |
| G109A      | 10          |
| G109A/L486A| 11          |
| G109A/I488A| 9           |

Surface expression levels were determined by flow cytometry using F-specific mAb F1a.

or 6-CF. Thus, these mutants do not cause either lipid mixing or content mixing. Previously, we determined that an increase in temperature can be a surrogate for HN triggering fusion. To examine temperature-dependent triggering of the F TM domain mutants, red blood cells (RBCs) were labeled with 6-CF and bound to effector CV-1 cells expressing HN and F proteins and labeled with SYTO-17 at 4 °C. The target–effector complexes were then incubated at 29, 37, or 42 °C for 15 min, and the number of fusion events was measured by confocal microscopy (Fig. 5c). For wt F, the number of dye transfer events increased with increasing temperature. The amount of fusion mediated by F TM domain mutants L486A and I488A increased between 29 and 37 °C, but increasing the temperature beyond 37 °C did not enhance fusion. The mutants L486A and I488A did not exhibit a hemifusion phenotype, i.e., transfer of the lipidic dye R18 to target cells in the absence of transfer of the aqueous dye 6-CF. It is possible that these mutations either stabilize the F protein and prevent F protein from attaining its lowest energy postfusion conformation or affect protein–membrane interactions that stabilize an intermediate in the fusion pathway. For example these mutant F proteins may be trapped at a folding/fusion intermediate that cannot be overcome by an increase in temperature.

F TM domain mutants L486A and I488A form the open-stalk and prehairpin intermediates of fusion

For PIV5 entry into cells by fusion, HN binds to its ligand, sialic acid, and by an unknown process begins the activation of F. The earliest stage of fusion that has been determined was inferred from functional properties of the N-1 peptide (which is derived from the HRA region of F). N-1 peptide can bind to the HRB region of F after HN has bound to target cells at 4 °C and inhibits fusion. Based on the atomic structure of prefusion F this step in fusion has been called the open-stalk conformation with the HRB helices melting and breaking the interactions at the base of the head but leaving the head domain largely intact. To determine if F TM domain mutants L486A and I488A can attain the F open-stalk conformation, a modified N-1 peptide, N-1-HAt, which contains an 11-residue HA tag (YPYDVPDYASL) at the C-terminus of N-1, was synthesized. Peptide binding was determined by the ability of the HA tag monoclonal antibody (mAb) to immunoprecipitate wt F protein. It was found that the N-1-HAt bound to F at 4 °C when target RBCs containing the HN receptor sialic acid were present but N-1-HAt did not bind to F as well in the absence of target RBCs. (Fig. 6a). To test if F TM domain mutants L486A and I488A reached the open-stalk stage of fusion, N-1-HAt peptide was incubated with cells expressing HN and one of the mutant F proteins at 4 °C in the presence of 0.5% hematocrit target RBCs (Fig. 6b). The HA tag mAb 12CA5 was used to immunoprecipitate the F protein containing bound peptide, and the total F in the lysate was immunoprecipitated with a polyclonal antibody (PAb) specific for F protein. N-1-HAt bound to wt F and F TM domain mutants L486A, and I488A, and the band for these mutants was generally more intense than that of the wt. This observation indicates the HRB region was accessible in all F proteins, and that the alanine substitutions in the TM domain might stabilize somewhat the formation of the open-stalk conformation of F protein.

The second known step in the F protein refolding event after formation of the open stalk intermediate is thought to be that HRA refolds and the fusion peptide is inserted into the target cell membrane to form the prehairpin intermediate. At this stage of fusion, the C-1 peptide, derived from the HRB region of F, specifically inhibits PIV5 F-mediated fusion by binding in the grooves on the outside of the HRA coiled coil. To capture the prehairpin conformation, the C-1 peptide was used in an RBC retention assay. To examine the F TM domain mutants for prehairpin formation, target RBCs labeled with 6-CF were bound at 4 °C to CV-1 cells that co-expressed F and HN and were labeled with SYTO 17. When the RBCs that have bound at 4 °C to the CV-1 cell are warmed to 37 °C, the majority of RBCs either fuse or are released due to the neuraminidase activity of HN that is active at 37 °C but not at 4 °C. In the presence of C-1 peptide, the F protein forms the prehairpin intermediate and the RBCs remain bound to the CV-1 cells because the fusion peptide has inserted into the RBC membrane, but further refolding is blocked and fusion is inhibited. Addition of C-1 peptide did not affect the number of RBCs bound at 4 °C (Fig. 6c). However, when cells expressing F TM domain mutants L486A and I488A
with bound RBCs were warmed to 37 °C in the absence of C-1 peptide, there was a near-complete loss of bound RBCs, suggesting that these mutants are not trapped at the prehairpin intermediate (Fig. 6c). To determine if these mutants reached the prehairpin intermediate or are unable to transition between the open-stalk conformation and the pre-hairpin intermediate, 40 μM C-1 peptide was added. At 37 °C there was an increase in the number of RBCs retained as compared to L486A and I488A at 37 °C without the addition of peptide, although the number was not as large as for wt F (Fig. 6c). These
data indicate that the F TM domain mutants form the prehairpin intermediate, but the reproducible lower amount of C-1 binding for mutants L486A and I488A could indicate some impairment in prehairpin formation for these two mutants.

**F TM domain mutants 486 and 488 have closely related postfusion conformations as wt F protein**

MAb 6–7 recognizes only the postfusion conformation of the F protein and not the prefusion conformation of the F protein. Thus, mAb 6–7 can be used to determine if F TM mutants L486A and I488A undergo the F protein refolding event that accompanies membrane fusion. It was found that although F TM domain mutants L486A and I488A cause greatly decreased fusion compared to wt F and TM domain mutants L486I and I488L, wt F and the four mutant F proteins exhibited the same mAb 6–7 reactivity at 4, 40, 43, 47, and 50 °C, as determined by flow cytometry, supporting the notion that the F protein TM domain
mutants can proceed through the known intermediates of fusion and that they have a conformation closely related to postfusion F, even though L486A and L488A are essentially fusion inactive (Fig. 6d). Thus, although the steady-state concentrations of the prehairpin intermediate might be affected by these mutations, the extent and rate of conversion to the postfusion form recognized by mAB 6-7 would appear to be essentially the same as that of wt.

The addition of a hyperfusogenic mutation can rescue fusion for F TM mutants L486A and I488A

It has been shown previously that a mutant of the W3A isolate of PIV5, S443P, demonstrates a lower temperature requirement for fusion activation, faster fusion kinetics, and independence of HN activation. Because for F S443P extensive syncytia formation occurs at room temperature in the absence of HN co-expression, F S443P and related mutants have been termed hyperfusogenic. It is thought that F mutation S443P destabilizes the interactions between the top of the HRB three helix bundle and the linker to the IgG-like domain II, hence lowering the energy barrier for conversion to the open-stalk conformation. Two other hyperfusogenic mutants that lower the temperature requirement for fusion and enable HN-independent fusion are Gly to Ala mutations in the fusion peptide G105A (previously referred to as G3A) and G109A (G7A). Both mutations are highly destabilizing: the G109A not only destabilizes the F protein on cell surface expression but also inactivates the F protein relatively quickly and inactivates F for fusion if target cells are not present.

It seemed possible that if F TM domain mutants are blocked in causing fusion at a stage beyond the prehairpin intermediate, then incorporation of a hyperfusogenic mutation into the TM domain mutants might overcome the fusion block, probably due to the increased kinetics of fusion that is thought to be caused by triggering a greater number of F trimers at any one time. The L486A and I488A mutations were introduced into three hyperfusogenic backgrounds, S443P, G105A, and G109A F, to create the double mutants S443P/L486A, S443P/I488A, G105A/L486A, and G109A/I488A (Fig. 7). The F double mutants were expressed in HeLa CD4 long terminal repeat (LTR) β-galactosidase (β-gal) cells and cell surface abundance was determined by flow cytometry. S443P/L486A, S443P/I488A, G105A/L486A, and G109A/I488A were surface expressed similarly to wt F, whereas the surface expression of G109A, G109A/L486A, and G109A/I488A was only 10% of that of wt F (Table 1). Expression in BHK-21F cells showed that F mutants S443P/L486A, S443P/I488A, G109A/L486A, and G109A/I488A exhibited extensive syncytia formation (hyperfusogenic) and syncytia formation was independent of HN expression (data not shown). F G105A/L486A and F G105A/I488A showed higher levels of syncytia formation than F L486A and F I488A but less than that of F G109A, consistent with earlier data for F mutant G105A. Thus, whereas F TM mutants L486A and I488A cannot cause fusion, the addition of the hyperfusogenic mutants to create double mutants lowers the energy barrier to fusion and overcomes the block to fusion caused by the L486A and I488A mutations. These data indicate that the F TM domain mutations L486A and I488A have not caused the F proteins to convert to an inactive “spent” conformation and the F TM mutant proteins have not veered off the fusion pathway. It may be that increasing the number of active F molecules increases the probability of a successful fusion event—perhaps the membrane
state and the number of active F molecules are parameters that interact in a manner unknown to determine a successful outcome. Alternatively, the “lifetime” of the prehairpin intermediate may be different between mutant and wt, and the presence of a greater number of active F molecules could enhance

![Graph](image-url)  
**Fig. 8.** Altering the curvature of the lipid membrane can rescue fusion for the F L486A and I488A mutants. (a) Quantification of cell–cell fusion in the dye transfer assay. Effector CV-1 cells were infected with vaccinia virus vTF7-3 and transfected with HN and wt F or mutant F DNA. RBCs were dually labeled with R18 (open bars) and the 6-CF (black bars). They were bound to CV-1 cells for 1 h at 4 °C and then incubated at 37 °C for 15 min before visualization by confocal microscopy. Prior to incubation at 37 °C, 10 μM LPC or 10 μM OA was added for 15 min at 4 °C. 10 μM LPC or 10 μM OA was also present when the temperature was raised to 37 °C. The means and error bars are from three microscopic fields. (b) Quantification of dye transfer with dually labeled RBCs as above, but with 0.5 mM CPZ was added for 1 min at room temperature and then washed out before the temperature was raised to 37 °C to trigger fusion.

![Graph](image-url)  
**Fig. 9.** Fusion activity of F proteins depends on the hydrophobicity of substituted residues at TM residues L486 and I488. (a) Cell–cell fusion of substitutions made at residues 486 and 488 were quantified using the luciferase reporter gene assay. Vero cells were cotransfected to express HN and wt F or mutant F proteins, and a luciferase reporter construct. At 16 h p.t. BSR T7/5 cells were overlaid on the Vero cells, and 6 h post overlay, the luciferase activity of each sample was read. Shown is the average of three experiments each done in triplicate and normalized to wt F. The next three panels show sequence suitability of protein sequences in the membrane quantified by the $E(z)$ scale. (b) The mean (continuous line) and one standard deviation above and below the average (dashed line) $E(z)$ score for 153 TM helices of known structure containing a minimum of 20 residues. Twenty-residue sliding windows were used for each helix sequence, with the residues renumbered such that the lowest-scoring (most membrane suitable) window occurs at residue number 0. The results for all 153 proteins were averaged after the renumbering. (c) $E(z)$ results for four viral TM proteins: PIV5 F, PIV5 HN, PIV5 SH, and influenza (flu) A M2. Horizontal lines show the mean (continuous) and one standard deviation above and below (dashed) values for the 153 TM helices in (a). (d) $E(z)$ results for four double mutants of PIV5 F. Residue positions L486 and I488 were mutated to alanine (A), threonine (T), tryptophan (W), and tyrosine (Y). Windows that do not include either mutation are shown in thick black (common region). (e) Atomistic model of the proposed extended TM region of PIV5 F. The canonical TM is shown in green, while the N-terminal (blue) and C-terminal (red) regions that are more suitable for the membrane than one standard deviation above the mean are also shown. This region is expected to be more suitable for the membrane than any window in 24 of the 153 TM domain sequences.
Fig. 9 (legend on previous page)
the probability of successful fusion. This second explanation is also consistent with the peptide pull-down results (Fig. 6).

Changing the curvature of the membrane rescues fusion of F TM domain mutants L486A and I488A

In addition to the F protein conformational intermediates of fusion, the changes occurring in the lipid bilayers can be broken down into several intermediates. It is thought that formation of the F protein 6-HB brings the membranes of the virus and the target cell together and exclusion of water molecules permits formation of the hemifusion stalk. This stalk is an initial lipid connection between the proximal membrane leaflets. The stalk then expands to form the hemifusion diaphragm, which is a single bilayer segment, before the fusion pore forms. Lipids can affect membrane fusion based on their molecular shape. The cone-shaped lipid stearoyl lysophosphatidylcholine (LPC) has a positive spontaneous curvature, which is predicted to hinder the transition to hemifusion when present in the outer bilayer. The cone-shaped lipid oleic acid (OA) has a negative spontaneous curvature, which favors hemifusion when present in the outer bilayer. The addition of LPC and OA to cells expressing HN and F proteins has been shown previously to inhibit and favor an increase in fusion, respectively. To examine whether addition of cone-shaped lipids would overcome the block in fusion of F TM domain mutants, LPC was added to CV-1 cells expressing HN and wt F, F L486A, or F I488A before and during binding of R18- and 6-CF-labeled RBCs at 4 °C. On warming to 37 °C, neither lipid mixing (R18 transfer) nor cytoplasmic content mixing (6-CF transfer) occurred between the RBCs and the CV-1 effector cells (Fig. 8a). In contrast, addition of OA to the effector cells not only increased wt F fusion, but also caused a significant increase in lipid and content fusion for F TM mutants L486A and I488A (Fig. 8a). Chlorpromazine (CPZ), a drug that causes positive curvature of the lipid bilayer, ruptures the hemifusion diaphragm and causes pore formation. The addition of CPZ to effector cells did not cause an increase in lipid or contents fusion for F TM domain mutants L486A or L488A (Fig. 8b). Thus, these data suggest that the block in fusion of F TM domain mutants L486A and I488A occurs during the lipid intermediate stages of fusion. It is not known at which stage of the lipid intermediates the F protein attains its final postfusion form, although it is usually considered that refolding of the prehairpin intermediate to the postfusion form occurs across the stages of the lipid intermediates. The augmenting of fusion by OA that confers negative curvature to the membrane suggests that the F TM domain mutants are delayed/arrested at the lipid stalk intermediate and do not reach the hemifusion diaphragm or fusion pore stages of fusion.

Evidence that the physical properties of the side chains of F TM domain residues 486 and 488 control fusion activity

If the TM region of F plays an active role in stabilizing the lipid stalk or other lipidic intermediates, then one might expect the nature of the residues near the head-group region to have a profound effect on fusion. Therefore, further substitutions were made at F TM domain residues 486 and 488 to test how the distinct potentials of different amino acid side chains to partition into different regions of the membrane would affect fusion activity. L486 and I488 were each substituted with I/L, V, F, C, A, G, T, W, or Y (Fig. 9). I, L, V, and F are residues that prefer to be buried deep in a bilayer, experiencing a favorable free energy of transfer from water to the bilayer center. By contrast, C, A, G, and T, show relatively flat potential profiles, indicating that transfer to the bilayer is neither strongly favored nor disfavored. Finally, W and Y have a strong tendency to localize in the head-group region of the bilayer, relative to either the bilayer center or the aqueous phase. The fusion activity of each of the mutants expressed in Vero cells was determined using a luciferase gene reporter assay. Fusion activity was severely decreased (less than 50% of wt F) for residue 486 or 488 substitutions A, G, T, W, and Y. Substitution for C yielded ~60% to 80% fusion, and substitutions for I/L, V, and F showed no difference in fusion activity compared to wt F protein (Fig. 9a).

When the fusion activity was plotted against the hydrophobicity index of each residue, those amino acid residues with a hydrophobicity index below 2 (A, G, T, W, and Y) show decreased fusion, whereas those amino acid residues with a higher hydrophobicity index exhibited wt F protein fusion activity (Supplemental Fig. 1). The E(2) potential was used to test further how the substitutions would affect the propensity of the TM sequence to insert into a bilayer. This potential, which is based on a statistical analysis of the crystallographic database of 153 membrane proteins, defines the potential for a given residue to reside at various depths in the membrane, and it is useful for determining which regions of a TM domain have the greatest preference to reside in specific regions of the bilayer.

Figure 9b shows a plot of the E(2) potential scores generated by computing the mean score for successive 20-residue segments across the sequence. The results are plotted for the average, and for segments with scores plus or minus a single standard deviation from the mean. In each case the sequences funnel down to a relatively sharp minimum. Because each point reflects the score for a 20-residue sequence, the minimum identifies the 20-residue segment with the highest propensity to lie vertical to the bilayer surface. This figure also compares the profiles of the F protein with that for the other two integral membrane proteins of the PIV5 (SH and HN) and the M2 proton channel of influenza A virus, the latter of which contains both a TM domain as well as a C-terminal cytoplasmic helical segment (Fig. 9c). The
sequences of SH, HN, and M2 funnel in the usual manner, and 20–30 residue stretches are observed with a high propensity to span the bilayer. Interestingly, however, the F protein sequence shows an unusually broad minimum, indicative of a much more extended sequence with some propensity to lie within the bilayer. This property is conserved among the paramyxovirus F proteins tested (measles virus, mumps virus, Newcastle disease virus and human parainfluenza viruses 2 and 3), although the profile for PIV5 F is the broadest (Supplemental Fig. 3). For PIV5 F, the absolute minimum in the curve suggests the TM helix begins at residue 485, in good agreement with the air-oxidation cross-linking data, showing this to be near the head-group region. However, with a very small energetic cost, the native TM helical bundle could translocate up or down through the membrane or lie at an oblique angle.

Interestingly, the scores for a 15-contiguous-point N-terminal minimum lies within one standard deviation of the mean value for the databank TM helices. This region includes a portion of the membrane-proximal end of the crystallographically defined structure of the HRB. Figure 9e illustrates the effects of the mutations on the cumulative $E(z)$ potential score for the N-terminal region for the mutations at residues 486 and 488. To help visualize the effects, the graphs were computed for double mutants, but the same trends hold for the individual mutants. The mutations change the 20-residue window of the protein with the most membrane-favorable sequence. Instead of the 20-residue window 485–504 being the most membrane favorable, the best window becomes 493–512. Additionally, with the exception of Trp, all the mutations have the effect of moving the most membrane suitable 20-residue window from 485–504 in the wild type to 493–512 in the mutants. Clearly, other regions of the protein are important for determining the boundary of the TM helix so it is difficult to interpret this as a literal movement of the TM in the resting state. Nevertheless, if helical movements are important for the fusion mechanism, these changes would upset the energetics of fusion. Figure 9e shows an atomistic model of the proposed extended TM domain region of PIV5 F. The canonical TM is shown in green, while the N-terminal (blue) and C-terminal (red) regions that are more suitable for the membrane than one standard deviation above the mean are also shown. The extended TM domain region is expected to be more suitable for the membrane than any window in 24 of the 153 TM domain sequences. 26

Discussion

For ion channel proteins that have multiple TM spanning domains, atomic structure determinations have enabled a distinction between those TM α-helices that serve architectural structural roles and those that act as the aqueous pore/ionic selectivity filter. 16 For the homotetrameric influenza virus M2 proton-selective ion channel protein, each polypeptide chain only spans the membrane once and the single hydrophobic domain found in each polypeptide chain has to act as the endoplasmic reticulum membrane insertion sequence, the membrane anchoring sequence and the pore and gate of the channel. 58,68,70–72 However, for the majority of integral membrane proteins that span a membrane once, the role of the TM domain besides being a membrane anchorage domain is largely unknown.

For many viral proteins that mediate membrane fusion, mutagenesis studies on the TM domain have yielded a wide variety of results. However, the preponderance of data indicates that the TM domain of these fusion proteins is not simply a string of hydrophobic amino acids that span a lipid bilayer but there is amino acid sequence specificity to the TM domain, implying specific roles of TM domain amino acid residues in fusion protein function. Specificity of amino acid residue implies specific structural features or interactions of these TM domain residues with the lipid bilayers and, thus, a direct role of amino acid side chains in the process of membrane fusion, further implying that membrane fusion is not simply a solely lipidic event.

To study the features of the PIV5 F protein TM domain for membrane fusion activity, alanine-scanning mutagenesis was used. The F protein TM domain was found to exhibit sequence dependence for fusion activity and it was also observed that a block of 20 leucine residues could not substitute for the entire PIV5 F TM domain even to obtain cell surface expression of F. Upon further substitution, residues L486 and I488 were found to play a key role in fusion, where altering the hydrophobicity of the side chain at these residues profoundly affected fusion activity (Fig. 9a). Although F mutants L486A and I488A were deficient for fusion, biological activity could be rescued by the addition of the destabilizing mutations F S443P, F G105A, or F G109A (Fig. 7), indicating F L486A and I488A were not trapped in an intermediate conformation nor had the mutant F proteins veered irreversibly off the fusion pathway. It is generally thought that F-mediated membrane fusion requires the action of several trimers and the hyperfusion phenotype of destabilizing mutants may be due to more synchronous F-activation events in an otherwise stochastic process. Thus, addition of a destabilizing mutation to F proteins that are blocked for fusion may simply alter the lifetimes of conformational intermediates, driving the fusion process by mass action. Detailed analysis indicated F L486A and F I488A form the prefusion conformation of F, the open-stalk intermediate, the prehairpin intermediate, and F can be converted to a conformation that is close to, or at the postfusion form (Fig. 1a–e). As discussed above, the precise timing of the conversion of the prehairpin conformation to the postfusion form with the formation of the lipid intermediates is not known, but refolding of F may occur across all the lipid intermediate stages. Evidence from studies of HIV
gp120/gp41 suggest some 6-HB formation occurs after a pore has formed.13,73

Thus, it seemed likely that the block in fusion for the F L486A and F I488A proteins corresponds to their inability to complete the lipidic stages of fusion (Fig. 1d and e). The data obtained using OA to confer negative spontaneous curvature of the membrane and concomitantly causing a large increase in fusion activity of F L486A and F I488A indicate that fusion is delayed/arrested at the lipid stalk intermediate (Fig. 1f–i). These findings suggest that the TM domain might play an active role in the lipidic steps in fusion, as discussed previously.16–19 In fusion, the formation of the 6-HB and the postfusion form of F brings together the TM domain and the FP, which excludes water between the membranes, and this local dehydration allows for membrane contact.2 There is a high-energy requirement for the presumptive next stage, the lipid stalk.74

One goal of this work was to define the structural features of the TM domain and to relate these to potential roles in the mechanism of fusion. The available atomic structures of ion channel proteins indicate that the multiple TM domains are mostly α-helical71 and for the very few known viral proteins, the structures of the TM domains of integral membrane proteins are α-helical.71,72,75 Even less is known as to whether the TM α-helices are together in the membrane forming helical bundles or whether each α-helix is separate from the others. Clearly, for the influenza virus M2 ion channel, the TM domains form a four-helix bundle.71,72 CryoEM studies on the human immunodeficiency virus fusion protein (Env) differ, as some reports indicate the TM domains of the fusion protein are apart, like the legs of a tripod,60,63 whereas other studies indicate the TM domains are together and form one helical bundle.32 The substitution of F TM domain residues with cysteine and cross-linking induced by air oxidation confirmed the N-terminal boundary of the TM in the cysteine and cross-linking induced by air oxidation allowing three consecutive residues to become flexible or are only partially folded, thereby allowing three consecutive residues to become very highly cross-linked (Fig. 3b). Quantification of the extent of disulfide bond formation and modeling studies indicate that the data fit best a model in which the F TM domain forms a 3-HB within the membrane. A break in the regular seven-residue period seen for coiled coils suggest that a residue was inserted near the center of the bilayer, resulting in either a pi bulge at residues 497–500 (Fig. 3d and e) or a “stutter” geometry (Fig. 3f). The PIV5 TM domain contains glycine at residues 494 and 497, but they do not appear to be essential to function or to mediate tight interhelical interactions. Residue L486 is predicted to be at the interface of the 3-HB, although the fusion data (in the presence of oleic acid) indicate this residue is likely to interact with the lipids of the bilayer. Presumably, the structural malleability that enables the residues close to the ectodomain to form disulfide bonds also enables residue L486 to populate different conformations that either face the interior of the predicted 3-HB or interact with the lipid bilayer.

The energy necessary to form the lipid stalk may be derived from the fusion protein, which may generate bilayer stresses that are relaxed by forming the stalk intermediate.2 For PIV5 F protein, the outer leaflet residues L486 and I488 may facilitate the negative curvature of the outer lipid leaflet that is necessary to merge the two bilayers, and residues with higher hydrophobicity may provide a more negative curvature (Fig. 1f–i).

The long potential length of the TM sequence of paramyxovirus F proteins (Fig. 9 and Supplemental Fig. 3) suggests an additional possible role for this domain in fusion. The length would allow the helix to move dynamically into different orientations and/or lie at extreme angles in the membrane. In this work, the extravalial end of the TM helix in the resting state was inferred from the Cys cross-linking as well as the E(z) potential, which were in good agreement. However, if this were the only orientation sampled by the helix during fusion, then mutation of residue I488 to Trp should not negatively impact function because it would be on the exposed face of the 3-HB located in the head-group region, where Trp is quite stabilizing. If, however, dynamic changes in the orientation within the bilayer were required, then this anchoring substituent would upset the energetic balance between the states adopted along the reaction coordinate. Given that the mutants appear to be defective in lipid stalk formation, and that the TM helix has a hydrophobic region both preceding and following it, we consider it possible that the TM region might act as a template to stabilize the lipid stalk intermediate. The N-terminal hydrophobic extension would encourage the diffusion of lipids out of the outer leaflet of the bilayer and along the stalk-like template, and initiating the formation of the lipid stalk (Fig. 10). Decreasing the hydrophobicity of the protein by mutating I488 or L486 to less polar residues or stabilizing the initial configuration by inclusion of Trp and Tyr would increase the energetic cost of translocation of the helix through the membrane and migration of lipids up the stalk.

Furthermore, the hydrophobic stretch at the cytoplasmic C-terminus of the TM helix might allow the TM 3-HB to translocate outward, extending the length of the exposed patch on the extracellular side of the bilayer (Fig. 10). Our calculations and experimental results suggest that the energy of the system is finely balanced, allowing this translocation to occur at minimal energetic cost. In the resting state, the TM bundle is stably inserted in the bilayer, but vertical translocations within the bilayer should occur at a relatively small energetic cost due to the lack of N-terminal anchoring residues and the hydrophobicity of the residues on both sides of the TM helix. More importantly, hairpin formation might provide a strong driving force for transloca-
tion of the TM helices in the bilayer. This conformational transition has the net effect of pulling the FP and the TM helix (located in opposing bilayers) toward one another, effectively applying a mechanical force favoring translocation of the N-terminus of the TM helix out of the bilayer, and the C-terminus of the TM helix deeper into the bilayer. The concomitant increase in length of the hydrophobic N-terminal extracellular region would provide a greater surface to serve as a template for nonbilayer lipidic intermediates. Finally, we note that in the prehairpin intermediate the 3-HB of HRA has a very hydrophobic surface prior to docking of the HRB helix and formation of the water-soluble postfusion 6-HB. Thus, the 3-HB of HRA could serve a similar role to encourage the initiation of a lipid-coated stalk projecting from the target cell’s membrane.

It is possible that as the last stage in the F protein refolding event the TM domain interacts with the FP and forms another 6-HB. The process of converting prehairpin F to the postfusion form of F may provide the necessary energy to exclude the water between the membranes and form the lipid intermediates (Fig. 1c–e). At the first stage of fusion, the outer leaflets of the two bilayers have just merged but have not mixed (hemifusion). The TM domain and FP may still be segregated in separate bilayers, and the interaction between the TM domain and FP could drive the formation of the hemifusion intermediate. Hyperfusogenic mutants G105A and G109A are located within the FP. These two mutants rescue fusion of L486A and I488A (Fig. 7). Although hypothesized previously that these destabilizing mutants G105A and G109A may cause hyperfusion by lowering the energy necessary to drive the various fusion intermediates, it is also possible that these FP mutants affect the potential interaction with the TM domain and destabilize and lower the energy requirement to mix the bilayers and drive the hemifusion intermediate and form the fusion pore (Fig. 1c–e).

In summary, the results presented here demonstrate that the TM helix of F plays an active role in fusion. It forms a well-defined TM bundle, and subtle modulations to its sequence give rise to large changes in activity. We further suggest that its unusual sequence characteristics might allow it to play an active role in templating the formation of the lipid stalk intermediate. We note that other proteins such as HIV gp41 also show unusually long hydrophobic TM helices, and changing its physical properties strongly impacts fusion. Additionally, dengue E protein (a class II fusion protein) has two TM helices, one of which has a long membrane

![Fig. 10. Hypothesized mechanism of lipid stalk intermediate stabilization. (a) Prefusion. A simplified representation of two PIV5 F protein trimers in the prefusion state is shown. The canonical TM domain is shown in orange. The regions hypothesized to enter the membrane in the lipid stalk intermediate are shown as white extensions of the TM domain. The fusion peptide is shown in red, HRA in green, HRB in blue, and the globular head is shown in yellow spheres. (b) Lipid stalk. As the lipid stalk forms, the extended TM domain completely enters the membrane environment, stabilizing the lipid stalk state and lowering the energetic cost of forming this intermediate. Arrows represent the movement of the globular head away from the stalk in the postfusion state. (c) Postfusion. Two F proteins are shown after refolding of the proteins into the postfusion state.](image-url)
suitable region that has an \( E(z) \) score one deviation from the average of 153 TM domains (Supplemental Fig. 4a). VSV (class III) has a single TM helix that also has a long membrane suitable region (Supplemental Fig. 4b). Although fusion is intrinsically a complex process with multiple transition states it would appear that viral fusogenic proteins share not only similar structures and conformational changes, but they might also specifically stabilize well-defined intermediates in the fusion pathway.

Materials and Methods

Cells, virus, and plasmids

BHK-21F, Vero, BSR T7/5, CV-1, and HeLa CD4 LTR \( \beta \)gal cells were grown in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS). BHK-21F cells were grown in DMEM supplemented with 10% tryptose phosphate broth, and HeLa CD4 LTR \( \beta \)gal cells were grown in DMEM supplemented with 200 \( \mu \)g/ml geneticin, 100 \( \mu \)g/ml hygromycin B, and 20 mM Heps (pH 7.4). The recombinant vaccinia virus (vTF7-3) that expresses T7 RNA polymerase was grown in CV-1 cells as described previously.53 pCAGGS plasmids encoding PIV5 G105A, or G107A F. Mutations were confirmed by nucleotide sequencing using an Applied Biosystems 3100-Avant automated DNA sequencer.

Expression of F and HN glycoproteins

PIV5 F and HN cDNAs cloned in the pCAGGS vector were expressed in BHK-21F, Vero, and HeLa CD4 LTR \( \beta \)gal cells by transient transfection using the Lipofectamine Plus expression system (Invitrogen, Carlsbad, CA) according to the manufacturer’s protocol. Transfected Vero cells were incubated for 4 h at 37 °C before the addition of DMEM containing 2% FBS and incubated a further 18 h at 37 °C. PIV5 F and HN cDNAs in the pGEM2X vector were expressed using the recombinant vaccinia virus–T7 RNA polymerase transient expression system (vacc T7).56 CV-1 cells in six-well dishes containing glass cover slides were infected at a multiplicity of infection of 10 plaque-forming units with vTF7-3 for 1 h at 37 °C. The cells were then transfected with 1.0 \( \mu \)g each of pGEM2X F and HN DNA using liposomes prepared as previously described.77 After 4 h at 37 °C, DMEM with 10% FBS was added and cells were incubated at 33 °C overnight.

Syncytia formation

Monolayers of BHK-21F cells in six-well plates were transfected with 1.0 \( \mu \)g each of pCAGGS PIV5 F and HN DNA as described above. At 20 h posttransfection (p.t.), cells were fixed and stained using a Hema 3 stain (Fisher Scientific, Pittsburgh, PA) according to the manufacturer’s instructions, and photographs were taken with a digital camera (DCS 760, Kodak, Rochester, NY) attached to an inverted phase-contrast microscope (Diaphot, Nikon, Melville, NY).

Luciferase reporter gene assay

To quantify cell–cell fusion, a luciferase reporter gene assay was performed as previously described.57 Briefly, Vero cell monolayers in six-well plates were transfected with 1.0 \( \mu \)g each of three plasmids, luciferase control DNA expressing the T7 promoter (Promega, Madison, WI), pCAGGS PIV5 F, and pCAGGS PIV HN. At 16 h p.t., BSR T7/5 cells expressing the T7 RNA polymerase were overlaid onto the Vero cells and incubated at 37 °C for 6 h. The monolayers were then washed, lysed, and clarified by centrifugation per the manufacturer’s instructions (Promega). For each sample, 150 \( \mu \)l of lysate was loaded into a 96-well plate. The luciferase activity of each lysate was quantified using 150 \( \mu \)l luciferase assay substrate (Promega) and an Lmax luminescence microplate reader (Molecular Devices, Sunnyvale, CA).

Dye transfer assays

Human RBCs were singly labeled with the aqueous dye 6-CF (Invitrogen) or dual labeled with 6-CF and the lipid probe octadecyl rhodamine F chloride (R18, Invitrogen). CV-1 cells were grown on glass cover slides and F and HN were expressed using the vacc T7 expression system. To visualize effector cells when using singly labeled RBCs, CV-1 cells were labeled with 1 \( \mu \)M SYTO-17 nucleic acid dye (Invitrogen) for 1 h at 37 °C. Analysis of lipid and aqueous dye transfer was performed as previously described.57 Fusion was quantified by counting positive cells by using a scanning confocal microscope (LSM 5 Pascal, Carl Zeiss MicroImaging, Inc., Thornwood, NY) and averaging the fusion events obtained from three separate fields. For \( \text{HNF}_{\text{indep}} \) independent retention of RBCs, SYTO-17 CV-1 cells were incubated with 0.1% hematocrit 6-CF-labeled RBCs. Following incubation with RBCs, cells were incubated at either 4 or 37 °C for 15 min. During this warming stage, 40 \( \mu \)M C-1 peptide was added to some of the samples. C-1 peptide was expressed in bacteria and purified as previously described.54 For the temperature dependence of dye transfer, samples were incubated at 29, 37, and 42 °C for 15 min after the binding of target RBCs. For dye transfer experiments using the addition of lipids, fresh solutions of 10 \( \mu \)M LPC (Avanti Polar Lipids, Birmingham, AL) or 10 \( \mu \)M OA (Sigma-Aldrich, St. Louis, MO) in phosphate-buffered saline (PBS) were made. After binding R18/6-CF dual-labeled RBCs as above, CV-1 cells were incubated in cold LPC and OA solutions for 15 min at 4 °C. The temperature was then shifted to 37 °C by changing the bathing solution with new PBS containing LPC or OA prewarmed to 37 °C and plates were incubated at 37 °C for 15 min as above. For dye transfer experiments with CPZ (Sigma), cells were prepared and R18/6-CF RBCs were bound as above. After raising the temperature to 37 °C for 15 min, 0.5 mM CPZ in PBS was added to the CV-1 cells for 1 min. Cells were extensively washed with PBS without drug.

Flow cytometry

To quantify cell surface expression and to determine the protein conformation of the F protein, monolayers of
HeLa CD4 LTR βgal cells in six-well plates were transfected with 1.0 μg of pCAGGS PIV5 F DNA as described above and flow cytometry was performed as previously described using fluorescein-isothiocyanate-labeled secondary antibody.59 For surface expression, mAb F15 was used at 1:100 dilution. To examine conformational rearrangements in the F protein, mAb 6–70 was used at 1:30 dilution. Before the addition of mAb 6–7, warmed PBS was added to the samples and the plates were incubated at 40, 43, 47, or 50 °C for 10 min and washed with cold PBS. The fluorescence intensity of 10,000 cells was measured by using a FACScalibur flow cytometer (Becton Dickinson, Franklin Lakes, NJ).

Capture of open-stalk intermediate

Monolayers of HeLa CD4 LTR βgal cells in 6-cm dishes were transfected with 2.0 μg each of pCAGGS PIV5 F and HN DNA as described above. At 18 h p.t., cells were starved with Cys- and Met-deficient DMEM for 30 min. The cells were labeled with 400 μCi of 35S-Promix (GE Healthcare Bio-Sciences, Piscataway, NJ) in 1 ml of Cys- and Met-deficient DMEM for 1 h. To allow for newly synthesized F proteins to reach the cell surface, the samples were chased with DMEM without serum for 2 h. The cells were washed with PBS and incubated with 1 ml of 0.5% hematocrit RBCs with or without 80 μl protein A-Sepharose beads overnight at 4 °C. The samples were washed three times with RIPA buffer containing 0.3 M NaCl, three times with RIPA buffer containing 0.15 M NaCl, once with 50 mM Tris buffer [0.25 mM EDTA (ethylenediaminetetraacetic acid), 0.15 M NaCl (pH 7.4)], and poly peptides were analyzed by SDS-PAGE on 15% acrylamide gels under nonreducing conditions in the absence of DTT.

Oxidative cross-linking

Monolayers of HeLa CD4 LTR βgal cells in six-well plates were transfected with 1.0 μg each of pCAGGS PIV5 F and HN DNA as described above. At 18 h p.t., cells were starved with Cys- and Met-deficient DMEM for 30 min. The cells were then labeled with 50 μCi of [35S]-Promix (GE Healthcare Bio-Sciences) in 1 ml of Cys- and Met-deficient DMEM for 1 h. The cells were then labeled with 50 μCi of [35S]-Promix (GE Healthcare Bio-Sciences) in 1 ml of Cys- and Met-deficient DMEM for 1 h. Cells were Dounce homogenized in cold RSB buffer [10 mM Tris (pH 7.4), 10 mM KCl, and 15 mM MgCl2], and 3 mM CuP (final concentration, freshly made) was added for 10 min at 37 or 4 °C. The reaction was stopped with 10 mM EDTA and 10 mM N-ethylmaleimide to chelate the copper and block free sulfhydryl groups. Samples were solubilized by adding 2× RIPA buffer plus 100 mM iodoacetamide and protease inhibitors as previously described81 and were clarified by centrifugation for 10 min at 55,000 rpm in a Beckman TLA100 rotor. Samples were centrifuged for 2 h at 4 °C with 10 μl of rabbit polyclonal anti-F2 peptide antiserum and then incubated with 40 μl protein A-Sepharose beads overnight at 4 °C. Samples were washed with RIPA buffer as above and poly peptides were analyzed by SDS-PAGE on 3.5% borate–acetate gels under nonreducing conditions in the absence of DTT.

Modeling of the TM domain

To better understand the periodic interface between the F protein TM domains, the raw radioactivity values were normalized by dividing the amount of disulfide formed by the total amount of protein (disulfide linked plus not linked). Because cross-linking is stronger toward the outside of the membrane and saturation can occur on the image plates of the Fuji Image Analyzer (Valhalla, NY), two sets of cross-linking data were combined for the analysis. First, cross-linking data using CuP at 37 °C were used for most of the TM positions, residues 492–509. To put the residues that are in the outer leaflet on the same scale, low-temperature (4 °C) CuP data were used for residue positions 485–491. Given the normalized data, a sine wave was fit according to the following formula:

$$y = \frac{a}{\frac{a}{b}} \times \sin((x + b)\pi/c) + d$$

where x is the residue number, y is the normalized degree of cross-linking, a is the amplitude of the sine wave, b is the phase offset, c is the α-helical periodicity, and d is the y offset of the sine wave. Values were fit by nonlinear regression. To visualize the cross-linking data on a single helix, cartoon representations of an ideal helix and a helix with a pi bulge were created with the interface residues labeled in spheres using the program PyMOL [DeLano, W. L. The PyMOL Molecular Graphics System (2002)].† The modeling protocol for the TM trimer is described in the Supplemental Methods.

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Supplementary Data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.jmb.2008.12.029

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