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The structural basis of paramyxovirus invasion

Charles J. Russell and Laura E. Luque

Department of Infectious Diseases, St Jude Children’s Research Hospital, Memphis, TN 38105-2794, USA

To deliver their genetic material into host cells, enveloped viruses have surface glycoproteins that actively cause the fusion of the viral and cellular membranes. Recently determined X-ray crystal structures of the paramyxovirus fusion (F) protein in its pre-fusion and post-fusion conformations reveal the dramatic structural transformation that this protein undergoes while causing membrane fusion. Conformational changes in key regions of the F protein suggest the mechanism by which the F protein is activated and refolds.

Paramyxovirus entry

The paramyxoviruses include many important human and animal pathogens such as measles virus, mumps virus, human respiratory syncytial virus (hRSV), human parainfluenza viruses 1–4 (hPIV1–4), Nipah virus, Hendra virus, parainfluenza virus 5 (PIV5, formerly known as SV5), Newcastle disease virus (NDV) and Sendai virus. To deliver their RNA genome into host cells, these enveloped viruses have evolved a membrane fusion machine that consists of two surface glycoproteins: a receptor binding protein and a fusion (F) protein. The receptor binding protein (called the HN, H or G protein depending on its biological properties) initiates membrane fusion by binding to receptors on the surface of host cells. The atomic structures of the hemagglutinin-neuraminidase (HN) proteins of NDV [1,2], hPIV3 [3] and PIV5 [4] have been determined in forms that are both bound and unbound to sialic acid receptor analogs. Although it remains unclear how the HN protein triggers the F protein to cause membrane fusion, the mechanism seems to involve changes in either the conformation or the oligomerization of the HN protein that transduces a signal to activate the F protein [5]. Once activated, the F protein initiates the process of membrane fusion.

The rearrangement of lipid bilayers during membrane fusion is thought to occur in a series of discrete steps that include dimpling, lipid stalk formation, hemifusion, transient pore formation and pore enlargement [6,7]. This process does not occur spontaneously; therefore, enveloped viruses have evolved surface proteins that undergo structural changes to do the work of membrane fusion. The paramyxovirus F protein is a member of the class I viral fusion proteins (Box 1). These proteins exist on the surfaces of infectious virions as trimers in metastable (high energy) conformations and become active for fusion usually only after proteolytic cleavage. For example, the paramyxovirus F protein is cleaved from an inactive precursor protein, F0, into the fusion-capable complex, F1 + F2. The class I viral fusion proteins also share similar domain structures that include a hydrophobic fusion peptide, two 4–3 heptad repeat regions (HRA and HRB) and a transmembrane domain (Figure 1a). Upon triggering, these proteins insert the fusion peptide into the host-cell membrane and refold HRA and HRB into a hairpin structure that brings the viral and cellular membranes together. X-ray crystal structures have recently been determined of the PIV5 F protein in the pre-fusion conformation [8] and the NDV and hPIV3 F proteins in the post-fusion conformation (Box 2) [9,10]. Along with the influenza virus hemagglutinin (HA) protein [11], the paramyxovirus F protein is currently the only other class I viral fusion protein for which atomic structures have been determined in both the metastable and hairpin conformations. Thus, the recently determined structures of the F protein provide fundamental insights into both paramyxovirus-mediated membrane fusion and the folding and refolding of metastable viral fusion proteins.

F protein structures

The PIV5 F0 protein in its pre-fusion conformation has a mushroom-like shape formed by a large globular head attached to a rod-like stalk (Figure 1b) [8]. The stalk is formed exclusively by HRB, which has a triple-stranded coiled-coil conformation. The base of the head is formed by two domains (DI and DII), which consist of residues that are predominantly located between HRA and HRB in the primary sequence. These domains are unique to the paramyxovirus F protein because the other class I viral fusion proteins have only short linker regions between HRA and HRB. In both the pre-fusion and post-fusion structures, DI forms a highly twisted β-barrel-like assembly and DII forms an immunoglobulin-like β-sandwich domain. Overall, DI and DII form a molecular scaffold that seems to help stabilize and then refold HRB and domain III (DIII).

The top of the head of the pre-fusion conformation of the F protein is formed by DIII, which consists of HRA, the fusion peptide and residues that are nearby in the primary sequence (Figure 1a). HRA is in a spring-loaded conformation at the top of the head, aimed towards the target membrane, and the fusion peptide is located on the side of the head, held in place by interactions with other residues in DIII and residues from DII. Such a configuration is reminiscent of the pre-fusion structure of the influenza virus HA protein [11]. However, HRA forms a helix–loop–helix structure in the HA protein and an 11-segment

Corresponding author: Russell, C.J. (charles.russell@stjude.org).
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structure consisting of α-helices, β-strands and turns in the F protein (Figure 1b).

The structures of the F proteins of NDV [9] and hPIV3 [10] in the post-fusion conformation reveal that DIII and HRB undergo dramatic rearrangements during membrane fusion. HRA refolds into a triple-stranded coiled coil that propels the fusion peptide ~ 115 Å from the side of the molecule into the target membrane. The HRB segments must also separate and then swing around the base of the head formed by DI and DII to bind to the HRA coiled coil and form the fusogenic hairpin structure [12,13] (Figure 1c).

Changes in F protein conformation during membrane fusion

The F protein is thought to adopt five distinct conformations during membrane fusion (Figure 2). The recently determined F protein structures represent the initial and final conformations of the F protein and suggest the mechanism by which it refolds into distinct intermediates. After cleavage of the F₀ precursor, the metastable conformation of the F protein is triggered to refold by binding of the HN protein to its receptor. The triggering of the F protein seems to involve a conformational change in HRB [14], and HRB has been shown to physically interact with the HN protein [15]. In the pre-fusion structure of the PIV5 F protein, residues at the nucleation site of the HRB coiled coil (residues 443, 447 and 449) interact with residues in the base of the F protein head. The importance of these HRB residues in the formation of an early F protein intermediate is confirmed by previous biochemical experiments, which show that mutations to residues 443, 447 and 449 alter the energy required to trigger the PIV5 F protein [16,17]. The next step in F protein refolding is the formation of a pre-hairpin intermediate in which HRA forms a triple-stranded coiled coil and the fusion peptide is inserted into the target membrane [14,17] (Figure 2d). Mutations to HRA residues 132 and 161 and fusion

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Box 1. Class I and class II viral fusion proteins [7]

- Class I viral fusion proteins include the paramyxovirus F protein, the influenza virus HA protein, the HIV gp160 protein, the Ebola virus GP protein and the severe acute respiratory syndrome (SARS) coronavirus S protein.
- Class I proteins cause membrane fusion by refolding into α-helical coiled-coil hairpin structures.
- Class II viral fusion proteins include the Dengue virus E protein, the West Nile virus E protein and the Semliki Forest virus E1 protein.
- The class II proteins of the flaviviruses and togaviruses cause membrane fusion by refolding their β-sheet-type domains into hairpin structures.

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Figure 1. Schematic representation of the paramyxovirus F protein. (a) Domain structure of the F protein. The locations of domain I (yellow), domain II (brown) and domain III (pink) are shown as solid lines above the diagram. The fusion peptide (FP, green), heptad repeat A (red), heptad repeat B (blue), transmembrane domain (TM, dotted line) and cytoplasmic tail (CT, dotted line) are also shown. (b) Model of the pre-fusion structure of the PIV5 F protein. The colors of the domains and regions are the same as in (a). HRA (red) is in a spring-loaded conformation and HRB (blue) forms the coiled-coil stalk. (c) Model of the post-fusion structure of the hPIV3 F protein. In the transition between the pre-fusion and post-fusion structures, domains I and II keep the same fold but reorient within the molecule. Domain III undergoes a dramatic refolding process, which leads to the formation of the coiled-coil hairpin structure that causes membrane fusion. In panels (b) and (c), α-helices are depicted as cylinders and β-strands as colored arrows.
peptide residues 105 and 109 increase the fusogenic activity of the PIV5 F protein [18–20]. The positioning of these residues at key structural points in the pre-fusion conformation of the F protein is consistent with mutations to these residues that destabilize the spring-loaded conformation and cause engagement of the F protein refolding process even in the absence of the HN protein trigger. The last conformational change in the F protein
during membrane fusion involves binding of the HRB segments into the grooves of the HRA coiled coil to generate a hairpin structure (Figure 2e). Because the fusion peptide is contiguous with HRA and the transmembrane domain is adjacent to HRB, the formation of the hairpin brings together the fusion peptide and transmembrane domain and, as a result, actively causes the fusion of the cellular and viral membranes [14].

Concluding remarks
High-resolution structures of the paramyxovirus F protein in its pre-fusion and post-fusion conformations now provide a structural basis for the analysis of the membrane fusion mechanism of the paramyxoviruses. In addition to explaining the phenotypes of previously characterized F protein mutants, these structures will enable future identification of residues that were previously unrecognized as important in regulating the progression of discrete refolding intermediates. A key unresolved issue in paramyxovirus fusion is the nature of the interaction between the HN protein and the F protein during fusion activation. Structures of both the HN protein and the F protein will certainly assist in the generation and testing of new hypotheses on the nature of this protein–protein interaction. The pre-fusion F protein structure will undoubtedly be used as a target in the structure-based design of fusion inhibitors. Peptides and small molecules that are designed to bind to the pre-hairpin intermediate of the F protein are potent inhibitors of virus replication [21–23]. It is possible that compounds that bind to the longer-lived native structure might have enhanced therapeutic properties. The successful use of the HIV fusion inhibitor enfuvirtide (T-20) in drug combination therapies is proof of the concept that fusion inhibitors can be clinically effective [24]. Although the metastable structure of the paramyxovirus F protein differs substantially from that of the influenza virus HA protein, key structural similarities such as the spring-loaded configuration of HRA and the positioning of the fusion peptide on the side of the molecule suggest that other class I viral fusion proteins such as HIV gp41 could also use similar strategies to stabilize and refold their pre-fusion conformations. Ultimately, further insights into how viruses invade host cells will depend on obtaining more structural information and performing additional biochemical experiments on the membrane fusion mechanisms of enveloped viruses.

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