The Many Mechanisms of Viral Membrane Fusion Proteins

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Abstract

Every enveloped virus fuses its membrane with a host cell membrane, thereby releasing its genome into the cytoplasm and initiating the viral replication cycle. In each case, one or a small set of viral surface transmembrane glycoproteins mediates fusion. Viral fusion proteins vary in their mode of activation and in structural class. These features combine to yield many different fusion mechanisms. Despite their differences, common principles for how fusion proteins function are emerging: In response to an activating trigger, the metastable fusion protein converts to an extended, in some cases rodlike structure, which inserts into the target membrane via its fusion peptide. A subsequent conformational change causes the fusion protein to fold back upon itself, thereby bringing its fusion peptide and its transmembrane domain—and their attached target and viral membranes—into intimate contact. Fusion ensues as the initial lipid stalk progresses through local hemifusion, and then opening and enlargement of a fusion pore. Here we review recent advances in our understanding of how fusion proteins are activated, how fusion proteins change conformation during fusion, and what is happening to the lipids during fusion. We also briefly discuss the therapeutic potential of fusion inhibitors in treating viral infections.

Keywords

Membrane fusion protein · Class I fusion protein · Class II fusion protein · Influenza HA · HIV Env · Low-pH activation · Receptor activation · Conformational changes · Membrane dynamics · Anti-fusion antivirals

1 Introduction

Fusion of enveloped viruses with host cells remains an important topic of research for two major reasons. First, it has recently become clear that fusion is a good target for therapeutic intervention (Kilby et al. 1998). Second, viral fusion reactions continue to serve as models for cellular fusion events. Although several viral fusion proteins, such as influenza hemagglutinin (HA) and the human immunodeficiency virus (HIV) envelope glycoprotein (Env), have emerged as paradigms, it is important to realize that there are many distinguishing features among viral fusion proteins (Table 1). Viral fusion proteins can be activated for fusion by different mechanisms. They have also been classified according to structural criteria. For some viruses, the viral receptor does not actively participate in fusion, whereas for others, one or more receptors are essential players. The location of the fusion peptide, critical for fusion, can vary. Finally, whereas some viruses require a single viral glycoprotein to mediate fusion, others require multiple viral glycoproteins. There are many excellent recent reviews on viral fusion and the glycoproteins that mediate this process (Durell et al. 1997; Eckert and Kim 2001; Heinz and
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Table 1. Viral membrane fusion proteins

| Family          | Viral proteins needed | pH of fusion | Class | Fusion peptide     |
|-----------------|-----------------------|--------------|-------|--------------------|
| Orthomyxovirus  | HA Low                | I N-terminal |       |                    |
| Alphavirus      | E1 Low                | II Internal  |       |                    |
| Flavivirus      | E Low                 | II Internal  |       |                    |
| Rhabdovirus     | G Low                 | Internal     |       |                    |
| Bunyavirus      | G1/G2 Low             | ? ?          |       |                    |
| Arenavirus      | GP Low                | ? ?          |       |                    |
| Filovirus       | GP Low<sup>a</sup>    | I Internal   |       |                    |
| Retrovirus      | Env Neutral<sup>b</sup> | I N-terminal, internal |       |                    |
| Paramyxovirus   | F,HN Neutral<sup>c</sup> | I N-terminal |       |                    |
| Herpesvirus     | gB, gD, gH, gL Neutral | ? ?        |       |                    |
| Coronavirus     | S Neutral             | I Internal   |       |                    |
| Poxvirus        | N.D. Neutral<sup>d</sup> | ? ?        |       |                    |
| Hepadnavirus    | S Neutral<sup>e</sup> | ? ?          |       |                    |
| Iridovirus      | N.D. N.D.            | ? ?          |       |                    |

<sup>a</sup> Inferred from infectivity assays.

<sup>b</sup> Most retroviruses fuse at neutral pH. MMTV appears to require low pH [Ross et al. (2002) PNAS 99:12386–90] to fuse. Avian retroviruses require receptor priming at neutral pH followed by exposure to low pH [Mothes et al. (2000) Cell 103:679–89; see text for a discussion of this model].

<sup>c</sup> Coronaviruses possess heptad repeats [Chambers et al. (1990) J Gen Virol 71:3075–80] characteristic of class I viral fusion proteins. Recent work indicates that they are, indeed, class I fusion proteins [Bosch et al. (2003) J Virol 77:8801–11].

<sup>d</sup> With infectivity assays, hepadnavirus uptake was shown to be pH-independent [Hagelstein et al. (1997) Virology 229:292–4]. However, recent studies have shown that duck hepatitis B virus may require low pH [Grgacic et al. (2000) J Virol 74:5116–22].

<sup>e</sup> The S protein contains a stretch of amino acids predicted to be a fusion peptide but has not been further characterized.

Allison 2001; Skehel and Wiley 2000; Weissenhorn et al. 1999). The goal of this review is to give the reader an appreciation for the diversity of viral fusion mechanisms.

2 Activation of Viral Fusion Proteins

All fusion proteins exist on virion surfaces in a metastable state in which the fusion peptide, a critical hydrophobic sequence, is hidden or shielded within the glycoprotein oligomer (Carr et al. 1997; Hernandez et al.
1996; Rey et al. 1995; Skehel and Wiley 2000; Wilson et al. 1981). After activation, the fusion peptides are rendered accessible for interaction with a target membrane. A major distinction among viral fusion proteins is the “trigger” for activation. There are two well-recognized mechanisms: (1) exposure to low pH and (2) specific interactions with target cell receptors at neutral pH. A third mechanism involving receptor priming at neutral pH followed by further activation at low pH was recently proposed (Mothes et al. 2000).

2.1 Low pH Activation

Orthomyxoviruses, togaviruses, flaviviruses, rhabdoviruses, bunyaviruses, arenaviruses, and, apparently, filoviruses require low pH to fuse with target membranes (Table 1) (Doms et al. 1985; Gaudin et al. 1999b; Stegmann et al. 1987; White and Helenius 1980). These viruses are endocytosed after binding to the target cell surface. The low-pH environment of the endosome activates the viral fusion protein to convert from a metastable state to one that is capable of driving fusion. Although the presence of a receptor may modulate the rate or extent of fusion (Ohuchi et al. 2002; Stegmann et al. 1996; White et al. 1982), receptors are not essential for low-pH-dependent fusion. Low-pH-dependent fusion generally occurs within seconds to minutes at 37°C but can also occur, albeit more slowly, at T<22°C.

Four main techniques have been used to assess whether a virus requires low pH to fuse. The first technique is testing the effects of agents, such as bafilomycin, that inhibit endosomal acidification. In some studies of this type, fusion has been measured directly by assessing the transfer of fluorescent probes from the virus to the target cell (Earp et al. 2003; Irurzun et al. 1997; Zarkik et al. 1997). In others, fusion has been inferred by monitoring postfusion events, such as the synthesis of viral DNA (Mothes et al. 2000).

A second test is to assess whether fusion of bound virions can be induced by briefly warming virus-cell complexes in low-pH medium (Mothes et al. 2000; White et al. 1980). A third test is to assess whether pretreatment of virions at low pH (in the absence of target membranes) inactivates the virus for fusion. Some (Bron et al. 1993; Corver et al. 2000; Di Simone and Buchmeier 1995; Korte et al. 1999; Nir et al. 1990; Stegmann et al. 1987), but not all (Puri et al. 1988), viruses that fuse at low pH can be inactivated by this method. Viral fusion proteins that are inactivated by low pH undergo irreversible conformational changes. In
the case of X:31 HA, this results in insertion of the fusion peptide into
the viral membrane (Korte et al. 1999; Weber et al. 1994).

The fourth test is to assess whether cells expressing the viral fusion
protein can fuse. Cell-cell fusion can be observed by light or fluores-
cence microscopy (Frey et al. 1995; Melikyan et al. 1997b; Mothes et al.
2000), or it can be scored with gene reporter assays that monitor inter-
actions of components from the fusing cells (Delos and White 2000;
Earp et al. 2003; Feng et al. 1996; Nussbaum et al. 1994). Although cell-
cell fusion assays are relatively simple to perform, the results do not
always correlate with virus-cell fusion or infection (Earp et al. 2003;
Lavillette et al. 1998; Schmid et al. 2000).

2.2
Receptor Activation at Neutral pH

Many enveloped viruses do not require low pH to fuse with target cells.
This has generally been established in controlled experiments using the
approaches described in Sect. 2.1. Viruses that can fuse at neutral pH in-
clude paramyxoviruses, herpesviruses, coronaviruses, poxviruses, and
most retroviruses (Table 1) (Hernandez et al. 1997; McClure et al. 1990;
Stein et al. 1987; Taguchi and Matsuyama 2002). The fusion proteins of
these viruses are activated via specific interactions with one or more re-
ceptors in the target cell membrane (Hernandez et al. 1996; Hunter 1997;
Stein et al. 1987). Viruses that can fuse at neutral pH are thought to do
so at the plasma membrane. However, they may also be able to fuse with
neutral-pH intracellular compartments (e.g., caveosomes) that can be
accessed through newly recognized endocytic pathways (Pelkmans and
Helenius 2003; Shin and Abraham 2001) (see also the chapter by
Sieczkarski and Whittaker, this volume). It is important to note, howev-
er, that viruses that can fuse at neutral pH may also possess the ability
to fuse at low pH (Earp et al. 2003; Fackler and Peterlin 2000). To date,
neutral-pH fusion has been found to display a sharp temperature thresh-
old, with little or no fusion occurring at T<20°C.

2.3
“Two-Step” Activation

Recently, a third model was proposed for the activation of alpharetro-
viruses. In this model, activation of the alpharetroviral Env begins with
receptor binding at neutral pH (at T>22°C) but is only complete after
exposure to low pH (Mothes et al. 2000). The role of low pH in this
“two-step” model is derived from two key observations: (1) The continuous presence of endosomal acidification inhibitors prevents production of alpharetroviral reverse transcripts, and (2) cells expressing Env and cells expressing the viral receptor only form large syncytia after exposure to low pH (Mothes et al. 2000). Our recent work indicates that alpharetrovirus fusion can proceed to the lipid mixing stage at neutral pH (Earp et al. 2003), and that receptor binding and low pH sequentially induce distinct conformational changes in the alpharetroviral Env (Matsuyama et al. 2004). Current work is now focused on determining the precise role of low pH in the fusion cascade.

3 Classification of Fusion Proteins Based on Structural Criteria

All viral fusion proteins contain a relatively large ectodomain, generally a single transmembrane domain, and all contain a cytoplasmic tail. So far, two major groups (class I and class II) have been defined based on structural criteria (Heinz and Allison 2001; Lescar et al. 2001) (Tables 1 and 2).

Class I fusion proteins are synthesized as precursors that are cleaved into two subunits by host cell proteases. In some cases (e.g., influenza HA), the two subunits remain associated through a disulfide bond; in others (e.g., HIV Env), the two subunits remain associated through non-covalent interactions. The proteolytic processing event that generates the two subunits is critical, as it creates the metastable state of the fusion protein (Chen et al. 1998). Class I fusion proteins exist as relatively long trimeric spikes in both their metastable and activated states. In their metastable states, they project perpendicularly to the viral membrane. The activated forms of the fusion subunits of known class I fusion proteins are highly α-helical (Skehel and Wiley 2000), and the final lowest-energy (which we will refer to as “postfusion”) forms (Fig. 1) contain “six-helix bundles” (Bullough et al. 1994; Carr and Kim 1993). All six-helix bundles contain a relatively long (65–115 Å) central N-terminal trimeric coiled-coil. Some (e.g., HIV Env, SIV Env, and paramyxovirus F) form six-helix bundles that extend to their membrane proximal ends [i.e., three C-terminal helices (Fig. 1A, green) pack in the grooves of the central coiled-coil (Fig. 1A, blue)]. Others display a mixture of helical and nonhelical segments that pack into the grooves of the central coiled-coil. For example, the HA2 subunit of influenza HA contains a relatively small six-helix bundle (Fig. 1, green/blue) at its membrane distal end,
followed by an extended chain (Fig. 1, yellow) that packs in the groove and extends to the N-terminal (membrane proximal) end of its central coiled-coil (Fig. 1B). Because of these variations, the postfusion forms of class I fusion proteins are often referred to as “trimers of hairpins” (Eckert and Kim 2001).

The general structure of class II fusion proteins is quite different from that of class I fusion proteins. A well-characterized example is the envelope glycoprotein (E) of tick-borne encephalitis (TBE) virus. During biosynthesis, TBE E and a second viral membrane glycoprotein, the precursor to the membrane protein (prM), form heterodimers. As virions ma-

### Table 2. Class I vs. class II viral membrane fusion proteins

| Property                                      | Class I                                      | Class II                                      |
|-----------------------------------------------|----------------------------------------------|-----------------------------------------------|
| Type of integral membrane protein            | Type I                                      | Type I<sup>a</sup>                           |
| Synthesized as                                | Inactive precursor                          | Inactive precursor<sup>b</sup>                |
| Exist on virion in                            | Metastable state                            | Metastable state                             |
| Orientation in virion (to membrane)           | Perpendicular                               | Parallel                                     |
| Converted to metastable state by             | Proteolytic processing within fusion protein precursor | Proteolytic processing of an associated protein |
| No. of subunits in fusion protein             | 2                                            | 1                                            |
| Major secondary structure of fusion subunit   | α-Helix<sup>c</sup>                          | β-Sheet                                      |
| Activated to fusogenic form by                | Low pH or cell receptor(s)<sup>d</sup>       | Low pH                                       |
| Oligomeric state of metastable protein        | Trimer                                       | Dimer                                        |
| Oligomeric state of fusion active protein     | Trimer                                       | Trimer                                        |
| Location of fusion peptide                   | N-terminal or internal                       | Internal loop                                |
| Structure of final fusogenic form             | Trimer of hairpins (coiled-coil)             | Trimer of hairpins (non-coiled-coil)          |

<sup>a</sup> The TBE E glycoprotein has two membrane anchoring segments near its C-terminal end [Heinz and Allison (2001) Curr Opin Microbiol 4:450–5].

<sup>b</sup> Known class II fusion proteins are activated by proteolytic cleavage of an accessory protein.

<sup>c</sup> The postfusion forms of all known class I fusion proteins are α-helical. The fusion subunit of metastable influenza HA is also highly α-helical, and this appears to be the case for a paramyxovirus F protein [Chen et al. (2001a) Structure 9:255–66]. Comparable information is not available for the metastable forms of other class I fusion proteins.

<sup>d</sup> In the case of paramyxoviruses, the receptor binding protein relays the information of receptor binding to the fusion subunit [Lamb 1993; Colman and Lawrence 2003]
ture, a host cell protease cleaves prM, resulting in reorganization of proteins on the viral surface (Allison et al. 1995). After prM cleavage, the E proteins exist as metastable homodimers. The ectodomains of the dimer are oriented antiparallel to one another. In further contrast to the trimeric class I fusion protein spikes, the ectodomains of the E homodimer lie parallel to the viral membrane and close to the surface. The TBE E protein is composed mostly of β-strand structure (Heinz and Allison 2001; Rey et al. 1995). The architecture of the Semliki Forest virus (SFV) spike, another well-characterized class II fusion protein, is similar to that of TBE E, but in this case, the metastable oligomer is a heterodimer of two membrane-anchored proteins, E1 and E2, with an associated small protein (E3).

4
Examples of Fusion Activation Mechanisms

In Sects. 4.1–4.4, we discuss a few examples of viral fusion proteins that employ different fusion mechanisms in more detail. These will include
examples of class I and class II fusion proteins, activated by low pH or by receptor interactions at neutral pH.

4.1 Influenza HA (Class I Fusion Protein, Low pH)

High-resolution structures are available for both the complete native (metastable) (Wilson et al. 1981) and activated (Bullough et al. 1994; Chen et al. 1999) forms of the influenza HA. On the viral surface, HA exists as a trimer of heterodimers (Fig. 2A). Each heterodimer consists of HA1, which contains the receptor binding domain (Fig. 2, gray), and HA2, which contains the fusion peptide (Fig. 2, red) and the transmembrane domain (located at the C-terminus). In the native (neutral pH) structure, the fusion peptide is buried within the HA oligomer. Three long helices, one from each monomer, come together to form the triple-stranded coiled-coil of the metastable trimer (Fig. 2A and B, blue and green).

Fig. 2A–D. Low-pH-induced conformational changes within influenza HA. HA1 is depicted in gray. The fusion peptide is red (HA2 residues 1–24). The coiled-coil is blue, with the C-terminal helix colored green. The C-terminal extended region is yellow. The transmembrane domain (not shown) attaches to the C-terminal end, indicated by “C”, of HA2. A model for conformational changes: A In the native, metastable, structure of HA, the fusion peptides are buried within the trimer interface. HA1 acts as a clamp to hold HA2 in a metastable state. HA2 is largely shielded by HA1. To illuminate the HA2 core, we have cartooned the portion of HA1 that covers HA2 as a simple (gray) line. B On exposure to low pH, the HA1 headgroups separate, allowing expulsion of the fusion peptide. C A loop-to-helix transition causes the fusion peptide to be repositioned to one end of HA2, where it can bind to the target membrane. D A helix-to-loop transition causes the C-terminal helix and the C-terminal extended region to reverse direction and bind to the grooves of the coiled-coil in an antiparallel orientation.
On exposure to low pH, HA undergoes dramatic conformational changes. The globular head domains separate, releasing the clamp that holds HA2 in its metastable state (Fig. 2B). As a result, the fusion peptide is exposed (Fig. 2C, red) at the top of an extended triple-stranded coiled-coil, in a position where it can interact with the target membrane. A helix-to-loop transition causes a short helix (Fig. 2D, green) and the C-terminal extended region (yellow) to flip up and run antiparallel to the central coiled-coil (Bullough et al. 1994). As a result, the fusion peptide and transmembrane domain are brought into close proximity at the same end of the molecule (Fig. 2D).

Many regions of HA are important for fusion. The fusion peptide is critical for hydrophobic attachment of the virus to the target membrane (Sect. 6.1). Mutations that prevent (1) globular head domain separation (Godley et al. 1992; Kemble et al. 1992), (2) the “B-loop”-to helix transition (Gruenke et al. 2002; Qiao et al. 1998), or (3) the C-terminal extended region from packing into the grooves of the final coiled-coil (Borrego-Diaz et al. 2003; Park et al. 2003) ablate the ability of HA to reach the lipid mixing stage of fusion. In our model (Gruenke et al. 2002), conversion of HA to a prehairpin intermediate (Fig. 2C) allows HA to bind to the target membrane. Further conversion to the hairpin structure (Fig. 2D) then drives the formation and opening of a fusion pore.

4.2 HIV Env (Class I Fusion Protein, Neutral pH)

Like influenza HA, HIV Env is synthesized as a single-chain precursor and cleaved during biosynthesis to yield gp120 and gp41. Native (metastable) HIV Env is a trimer of the heterodimers of gp120 (the receptor binding subunit) and gp41 (the fusion subunit). Env is activated for fusion (at neutral pH) after sequential binding to CD4 and a coreceptor (a chemokine receptor). Binding of Env to CD4 causes conformational changes in Env that permit binding to the coreceptor. After coreceptor binding, additional conformational changes occur in Env that lead to fusion (Eckert and Kim 2001).

Crystal structures exist for the core of the gp120 subunit (Kwong et al. 1998) as well as for the postfusion (Fig. 3, Step 6) form of gp41 (Chan et al. 1997; Weissenhorn et al. 1997). However, there is not yet a crystal structure of the native (metastable) Env trimer. Therefore, a detailed picture of HIV Env activation via receptor interaction is not available. We presume that the first steps of Env activation are separation of the
globular head domains, expulsion of the fusion peptide, and extension of gp41 into a prehairpin intermediate (Fig. 3, Step 1). Several lines of evidence indicate the existence of the prehairpin intermediate. For example, peptide analogs of the C-terminal helix (Fig. 3, green) strongly inhibit HIV fusion and infection (Chan and Kim 1998; Kilby et al. 1998). Also, a synthetic peptide corresponding to the C-terminal helix coimmunoprecipitates with HIV Env after engagement of receptors (Furuta et al. 1998; He et al. 2003). The C-terminal helix then packs, in an antiparallel fashion, into the groove of the N-terminal coiled-coil (Fig. 3, Step 5). Because the C-terminal helices of gp41 extend along the entire length of the N-terminal coiled-coil, this packing would bring the fusion peptide and transmembrane domain very close together. The transition to the six-helix bundle drives membrane merger (Melikyan et al. 2000a). Moreover, complete six-helix bundles are needed to form “robust” fusion pores (Markosyan et al. 2003).

As mentioned above, HIV studies, primarily using epitope accessibility assays, have indicated that engagement of HIV receptors induces con-
formational changes in gp120 and gp41 (Eckert and Kim 2001; Xiang et al. 2002). A remaining issue for all receptor-activated viral fusion proteins is how information is transmitted (after receptor binding) through the receptor binding subunit to the fusion subunit. Such transmission is essential to allow rearrangements in the fusion subunit (e.g., six-helix bundle formation) that drive fusion. For HIV, part of the mechanism may involve reduction of one or more disulfide bonds in gp120 (Abrahamyan et al. 2003; Barbouche et al. 2003; Fenouillet et al. 2001; Gallina et al. 2002). In murine retroviral Envs, a proline-rich hinge region appears to relay receptor binding information from the N-terminal to the C-terminal region of the receptor binding subunits (SU) (Barnett and Cunningham 2001; Lavillette et al. 2001). Because the proline-rich region of SU is linked to TM by a disulfide bond (Pinter et al. 1997), this may provide a relay system to trigger conformational changes in the fusion subunit. Clearly, the molecular pathways by which receptor-activated fusion proteins change from their metastable to their activated forms need to be defined.

In Fig. 3, we show a working model for HIV Env-mediated fusion. It is derived in part from studies with influenza HA, and it is similar to other HIV fusion models (Eckert and Kim 2001). Our hypothesis is that all class I fusion proteins will employ similar mechanisms. We note, however, that even in the case of influenza HA, alternate models are still entertained (see Fig. 2 in Jahn et al. 2003). Furthermore, others have suggested that different class I fusion proteins may use fundamentally different mechanisms (Chen et al. 2001a).

The features that we predict will be common to the fusion mechanisms of all class I fusion proteins (Fig. 3) include: (1) conversion from a metastable state to an activated state, (2) exposure and repositioning of the fusion peptide for binding to the target bilayer, (3) recruitment of several activated fusion proteins to a fusion site (Blumenthal et al. 1996; Danielli et al. 1996; Markovic et al. 2001; Markovic et al. 1998), and (4) subsequent conformational changes that result in close apposition of the fusion peptide and the transmembrane domain.

4.3 Paramyxovirus F Proteins (Class I Fusion Protein, Neutral pH, Attachment Protein Assisted)

The viral fusion proteins that have thus far been discussed in detail contain a receptor binding domain (e.g., the gp120 subunit of HIV Env) within the fusion protein spike. In other cases, the receptor binding do-
main resides in a separate viral spike. Paramyxoviruses have an attachment protein spike and a separate fusion (F) protein spike. Most, but not all, paramyxoviruses require both the attachment protein and the F protein for fusion (Bagai and Lamb 1995; Paterson et al. 2000). In most cases, the attachment protein must come from the same paramyxovirus as the fusion protein (Bossart et al. 2002). In the few cases in which the F protein is sufficient, fusion is enhanced if the attachment protein is also expressed (Bagai and Lamb 1995). The need for the attachment protein can be overcome by mutations in the F protein (Paterson et al. 2000; Seth et al. 2003) or by conducting fusion reactions at T>37°C (Paterson et al. 2000; Wharton et al. 2000). Paramyxovirus fusion proteins thus represent special cases of receptor-activated fusion proteins, in which receptor activation is communicated from one viral spike glycoprotein to another.

F proteins are proteolytically cleaved during biosynthesis to generate two disulfide-bonded subunits, F₁ and F₂ (Begona Ruiz-Arguello et al. 2002; Gonzalez-Reyes et al. 2001; Lamb 1993), found as metastable trimers of dimers (Baker et al. 1999) on virions. It has been suggested that binding of the attachment protein to a host cell receptor causes conformational changes in this protein, which in turn cause activating conformational changes in the metastable F protein (Lamb 1993; Russell et al. 2001; Takimoto et al. 2002). The exact mechanism by which attachment proteins activate F proteins is not known, but several groups have provided evidence for cross talk between attachment and F proteins (Bossart et al. 2002; Deng et al. 1999; McGinnes et al. 2002; Stone-Hulslander and Morrison 1997; Takimoto et al. 2002; Yao et al. 1997).

The post-fusion form of the F protein from the paramyxovirus SV5 contains a six-helix bundle (Baker et al. 1999). Similar to HIV Env (He et al. 2003; Kilby et al. 1998; Munoz-Barroso et al. 1998) and other retroviral fusion proteins (Earp et al. 2003; Netter 2002), peptide analogs of the N- and C-terminal helices of paramyxovirus six-helix bundles are potent inhibitors of fusion and infection (Bossart et al. 2002; Joshi et al. 1998; Lambert et al. 1996; Young et al. 1999). As is also the case for HIV Env (Markosyan et al. 2003; Melikyan et al. 2000a), a recent study showed that conversion of the SV5 F protein to a six-helix bundle drives membrane fusion (Russell et al. 2001).

Issues yet to be addressed for paramyxoviruses are the structure of the complete native (metastable) F trimer and how it is converted to its activated form. The first glimpses at the metastable and postfusion states of the F trimer came from EM observations of the respiratory syncytial virus (RSV) F protein. Preparations of purified recombinant F protein
contained both cone-shaped rods and “lollipop”-shaped structures. On storage, there appeared to be a shift from the cone-shaped to the “lollipop”-shaped structures (Calder et al. 2000). Examination of F complexed with specific monoclonal antibodies suggested that the “lollipop” structures contained six-helix bundles composed of N- and C-terminal heptad repeats (Calder et al. 2000).

A high-resolution structure of an F protein ectodomain was recently presented (Chen et al. 2001a). The protein used for the analysis contained a mixture of precursor F0 and proteolytically cleaved F. It also apparently lacked the second heptad repeat, which forms the C-helix in the postfusion form. This trimeric F protein structure is fundamentally different from that of influenza HA; the N-terminal end of its coiled-coil is positioned near the viral membrane end of the molecule (i.e., opposite the orientation of the coiled-coil in the metastable HA trimer). If this F protein structure represents the native metastable F trimer, then it suggests a mechanism of fusion activation for F fundamentally different from that for HA (Chen et al. 2001a). Additional work is needed to test this idea.

4.4 TBE E and SFV E1 (Class II Fusion Proteins, Low pH)

All known class II fusion proteins are activated by low pH. However, the mechanism by which class II fusion proteins are initially activated is quite different than the mechanism by which class I fusion proteins are initially activated. For example, the ectodomain of the TBE glycoprotein forms an antiparallel dimer that lies parallel and close to the viral membrane (Fig. 4B). At low pH, the TBE E homodimer converts to an E homotrimer (Allison et al. 1995; Heinz and Allison 2001; Stiasny et al. 2001). This transformation is thought to occur in two steps: dissociation of the E homodimer, followed by reassociation of E trimers (Stiasny et al. 1996). Membrane binding occurs after dimer dissociation and promotes the formation of E homotrimers (Stiasny et al. 2002). Homotrimer formation may involve interactions between α-helices in the stem region of the E protein (Allison et al. 1999).

The SFV fusion protein also converts from a dimer to a trimer during fusion activation. On native virions, E1 exists as a tight heterodimeric complex with a second membrane protein, E2. On exposure to low pH, E1 dissociates from E2, changes conformation, and forms highly stable E1 homotrimers (Ahn et al. 1999; Kielian 1995; Wahlberg et al. 1992; Wahlberg and Garoff 1992). During this process, E1 binds hydrophobically through its fusion peptide to target membranes and mediates fu-
Similar to TBE E, it appears that binding to the target bilayer fosters formation of the activated E1 homotrimer (Kielian 1995; Kielian et al. 2000); the fusion peptide and the transmembrane domain of E1 appear to be important for E1 homotrimer formation (Kielian et al. 1996, 2000; Sjoberg and Garoff 2003). Thus both class I and class II viral fusion proteins appear to function as trimers during fusion. It has been proposed that activated TBE E (Helenius 1995) and SFV E1 stand up as trimeric spikes and present their fusion peptides to the target membrane (Fig. 4B, right). This would be analogous to Fig. 3, Step 1. If this occurs,
then the spike would have to refold to bring the viral and cellular membranes together (e.g., analogous to Steps 4 and 5 in Fig. 3). Very recent evidence indicates that this is, indeed, the case (Bressanelli et al. 2004; Gibbons et al. 2004; Modis et al. 2004).

5 Membrane Dynamics During Fusion

Thus far we have focused on the conditions that elicit viral fusion reactions and the conformational changes in viral fusion proteins necessary for fusion. However, it is the viral and cellular bilayer membranes that merge during fusion. Lipid bilayers are stable structures that do not fuse spontaneously. Fusion proteins have evolved to catalyze the necessary lipid rearrangements. We now review a lipid rearrangement model and focus on the roles of different regions of viral fusion proteins in choreographing the structural changes that the membranes undergo throughout the fusion cascade (Fig. 3).

The favored model for the lipid transition state during membrane fusion is the stalk model. In this model, two opposing membranes bend toward each other, creating “dimples” (when viewed from the trans surface) or “nipples” (when viewed from the cis surface) (Fig. 3, Step 4). Nipples continue to bend until they meet. The two cis leaflets then merge, creating a lipid stalk (see Fig. 2 in Kozlovsky and Kozlov 2002) that proceeds to a state of local hemifusion (Fig. 3, Step 5). In a second step, transient fusion pores form, which give rise to stable pores (Fig. 3, Step 6).

The first direct visualization of a lipid stalk intermediate was achieved by electron diffraction studies of the effect of sequential dehydration on lipid bilayers composed of a lipid that has negative spontaneous curvature (Yang and Huang 2002). The stalk intermediate was stable at intermediate relative humidities. The results suggested that both the formation of a lipid stalk and its transition to a conformation that can be equated with pore formation require external forces.

Cellular membranes do not have spontaneous negative curvature and are highly hydrated. Membrane curvature can be promoted by introducing defects into the contacting bilayers. Thus roles for the fusion protein include pulling the fusing bilayers toward one another (dimpling), dehydrating the membranes, and creating membrane defects that lower the energy barrier for stalk and pore formation. Two intermediates in HIV fusion have been trapped: one in which the two membranes are joined
by activated Envs, but are not yet fused (Melikyan et al. 2000a), and one in which small, “labile” pores have formed that can either expand into stable, “robust” pores or return to the prefusion state (Markosyan et al. 2003). These observations suggest a role for the fusion protein in formation and stabilization of both the fusion stalk and the fusion pore.

The mechanism by which a small pore enlarges is not known. However, several possibilities have been proposed. One is that the initial fusion pore is formed by a small number of activated fusion proteins. Additional activated fusion proteins then move into the fusion site to buttress and stabilize the pore, thereby allowing it to expand (Kozlov and Chernomordik 2002). Another possibility is that multiple small fusion pores coalesce to form larger ones. This was supported by EM visualization of HA-mediated fusion, in which multiple dimples/nipples were arranged circularly and lipid fragments were seen at the center of a fusion ring (Kanaseki et al. 1997).

6 Membrane-Interacting Regions of Viral Fusion Proteins

As discussed above, roles for the fusion protein in the fusion cascade (Fig. 3) include pulling the fusing bilayers toward one another (dimpling) and creating membrane defects that lower the energy barriers for stalk formation and fusion pore opening/enlargement. The fusion peptide and the transmembrane domain must remain stably associated with the target and viral membranes, respectively, for fusion to occur. Once the fusion peptide is stably associated with the target bilayer (Fig. 3, Step 2), we envision that rearrangements in the fusion protein ectodomain that bring the fusion peptide and transmembrane domains close together (Fig. 3, Step 4) result in dimpling of membranes toward one another. In addition to serving as critical membrane anchors, the fusion peptide and the transmembrane domain likely create membrane defects that facilitate the next stages of fusion. Here, we review information about the structure and function of the fusion peptide and the transmembrane domain during fusion. We also review evidence that juxtamembrane sequences, on both sides of the transmembrane domain, participate in fusion.
6.1 The Fusion Peptide

Fusion peptides are relatively apolar sequences that interact with membranes and are central to viral fusion reactions (Martin and Ruysschaert 2000; Martin et al. 1999; Skehel et al. 2001; White 1990). They have been

| N-terminal | Internal |
|------------|----------|
| **Class I** | **Ebola GP:** | GAAIGLAWIPYPGPAA |
| Influenza HA2: | ASLV gp37: | IFASILAPGVAAQAL |
| Sendai F1: | **Class I** | DYQCKVYTGVYPFMWGGAYCFCD |
| Resp. Syn. F1: | TBE E: | DRGWGNHCGLFGKGSIVA |
| HIV gp41: | unclassified | VSV G: QGTWLNPFPQSCGYATV |

**Fig. 5A, B.** Characteristics of viral fusion peptides. A Selected viral fusion peptide sequences. N-terminal (Skehel et al. 2001) and internal (Delos et al. 2000) fusion peptide sequences are aligned according to their first noncharged residue. B Model of HA fusion peptide structure in target membrane at pH 5 (adapted from Tamm et al. 2002). The fusion peptide (red) resides in the target membrane in a kinked structure composed of two α-helices, each penetrating the outer leaflet. The glycine ridge is depicted by a yellow box, the hydrophobic interior face by cyan ovals, and the surface charged residues by blue squares. “C” denotes the direction of the HA2 ectodomain.
classified as N-terminal or internal depending on their location within the fusion subunit (Table 1). Although fusion peptides are highly conserved within each virus family, there is little sequence similarity between fusion peptides of different families (Fig. 5A). Generally, however, fusion peptides contain a high percentage of glycines and/or alanines, as well as several critical bulky hydrophobic residues (Martin and Ruysschaert 2000; Martin et al. 1999; Skehel et al. 2001; Tamm and Han 2000; Tamm et al. 2002).

6.1.1 Structure of N-terminal Fusion Peptides

A significant body of work has emerged on the structure and function of synthetic fusion peptides (Martin and Ruysschaert 2000; Martin et al. 1999; Skehel et al. 2001; Tamm and Han 2000; Tamm et al. 2002). Synthetic fusion peptides are disordered in solution but ordered (α-helix and/or β-sheet) when they associate with membranes. The N-terminal fusion peptides that have been studied insert into membranes at oblique angles and do not penetrate the inner leaflet of the membrane. In general, mutations that abrogate fusion reduce the ability of synthetic fusion peptides to insert at oblique angles and to disrupt membranes (Martin et al. 1999). Contradictory conclusions on the precise structure of synthetic fusion peptides in membranes likely stem from the general low solubility of the peptides in aqueous solution and the different experimental methods employed (Tamm et al. 2002).

To circumvent solubility problems, a polar sequence was added to the C-terminal end of the influenza HA fusion peptide, rendering it soluble in both aqueous and hydrophobic environments (Han et al. 2001). At pH 5, the HA fusion peptide consists of an N-terminal helix, a kink, and a short C-terminal helix (Fig. 5B). Both the N- and C-terminal helices penetrate the outer leaflet of the target bilayer. The kink remains at the phospholipid surface; the interior (lipid-facing surface) of the kink is lined with hydrophobic residues. The conserved glycines form a ridge along the outer face of the N-terminal helix. Three charged residues are also found on the outer face (Fig. 5B). An HA in which the conserved glycine at the beginning of the fusion peptide (Gly1) has been changed to valine cannot mediate fusion. If Gly1 is changed to serine, HA mediates only hemifusion or only forms small nonexpanding fusion pores (Qiao et al. 1999; Skehel et al. 2001). Interestingly, these mutant fusion peptides have membrane-associated structures and orientations significantly different from those of the wild-type fusion peptide (Li et al.
Simulations suggested similar membrane penetrating orientations for the HIV fusion peptide and two fusion-defective mutants (Kamath and Wong 2002).

### 6.1.2 Structure of Internal Fusion Peptides

In addition to a significant number of apolar residues, many internal fusion peptides contain a conserved proline at or near their centers (Fig. 5A). Mutagenesis of this proline in the avian sarcoma/leukosis virus (ASLV) EnvA fusion peptide suggested that it stabilizes a \( \beta \)-turn (Delos et al. 2000). This, coupled with the observation that mutating two cysteines that flank the fusion peptide abolishes fusion activity (Delos and White 2000), suggested that the internal EnvA fusion peptide exists as a looped structure stabilized by a disulfide bond. The ability of the Ebola virus fusion protein, which also contains an internal fusion peptide, to support infection was similarly inhibited when its central proline and flanking cysteines were mutated (Ito et al. 1999; Jeffers et al. 2002). A similar mutation of a proline within the predicted turn segment of the candidate fusion peptide of VSV G also significantly decreased fusion and abolished infectivity (Fredericksen and Whitt 1995). The idea of loop structures for internal fusion peptides is further supported by the known looped structure of the TBE E and SFV EI fusion peptides (Rey et al. 1995; Allison et al. 2001; Lescar et al. 2001). In some cases, two or more noncontiguous sequence loops may function as a collective fusion peptide (Gaudin et al. 1999a; Li et al. 1993).

Like N-terminal fusion peptides, internal fusion peptides contain a significant number of glycines and hydrophobic residues (Fig. 5A). Changing either of two glycines within the SFV EI fusion peptide to alanines altered the pH threshold for fusion, and changing one of the glycines to aspartic acid abolished fusion (Duffus et al. 1995; Kielian et al. 1996). Alteration of hydrophobic residues at the beginning, middle, or end of the (internal) ASLV EnvA fusion peptide to charged residues impaired the ability of EnvA to mediate fusion (Hernandez and White 1998). Similarly, a tryptophan and a glycine are critical for Ebola GP-mediated infection (Ito et al. 1999). Also, a bulky hydrophobic residue is needed at the tip of the TBE E fusion peptide loop (Rey et al. 1995) (Fig. 4A, red). Collectively, these results suggest that internal fusion peptides function as loops that require a mixture of hydrophobic and flexible residues, similar to those found in N-terminal fusion peptides.
6.1.3
Roles of Fusion Peptides

Fusion peptides appear to act at several steps along the fusion pathway. As demonstrated by mutants in which apolar fusion peptide residues were changed to charged residues (Freed et al. 1992; Gething et al. 1986; Hernandez and White 1998; Schoch and Blumenthal 1993), fusion peptides clearly play an important role in anchoring the fusion protein to the target membrane (Fig. 3, Step 2). The energy provided by inserting the fusion peptides of a single HA trimer into a membrane would be sufficient to initiate stalk formation (Gunther-Ausborn et al. 2000). The fusion peptide may also assist in creating the stalk by displacing water from the lipid-water interface, thus decreasing the repulsive force between the two fusing membranes (Tamm and Han 2000). Fusion peptides may also function in fusion pore opening. In support of this possibility is the observation that an HA mutant in which Gly1 was changed to serine mediates extensive lipid, but not content mixing (Qiao et al. 1999). Furthermore, defects in syncytium formation and infectivity were observed for HIV Env harboring the mutation V2E in its fusion peptide (Freed et al. 1992). Biophysical studies comparing a synthetic fusion peptide harboring this mutation with the wild-type peptide suggested a requirement for fusion peptide aggregation in the creation of the HIV fusion pore (Kliger et al. 1997; Pereira et al. 1995).

6.2
The Transmembrane Domain

Studies with chimeric fusion proteins have suggested that the transmembrane domains of some viral fusion proteins do not require a specific sequence to support fusion (Armstrong et al. 2000 and references therein). In contrast, studies with glycosylphosphatidylinositol (GPI)-anchored fusion proteins have demonstrated that there is a strict requirement for a proteinaceous membrane anchor for fusion proteins to efficiently mediate the transition from hemifusion to full fusion (Kemble et al. 1994; Melikyan et al. 1997a; Tong and Compans 2000). There also appears to be a minimum length for the fusion protein transmembrane domain to be able to support this transition (Armstrong et al. 2000; West et al. 2001). Therefore, it has been suggested that fusion protein transmembrane domains must span both leaflets of the viral bilayer to mediate fusion pore opening (Armstrong et al. 2000).
The transmembrane domains of some fusion proteins appear to have specific amino acid requirements for fusion function. For example, a conserved positively charged residue in the middle of the transmembrane domains of certain retroviral Envs appears to be important for the ability to mediate fusion and infection (Einfeld and Hunter 1994; Owens et al. 1994; Pietschmann et al. 2000; West et al. 2001). Two glycine residues in the transmembrane domain of VSV-G appear to be important for the transition from hemifusion to full fusion (Cleverley and Lenard 1998). Studies using a synthetic peptide corresponding to the mutant VSV-G transmembrane domain (Dennison et al. 2002) suggested that the VSV-G transmembrane domain lowers the energy barrier for fusion and stabilizes the transient fusion pore, thereby promoting its conversion to a stable fusion pore. Two glycines may allow the VSV-G transmembrane domain to adopt alternative conformations under different conditions, and such flexibility may be important for function. The transmembrane domain of HA from the Japan (Melikyan et al. 2000b), but not the X:31 (Armstrong et al. 2000), strain of influenza appears to require a glycine near the middle. An ability to adopt alternative conformations was also invoked to explain the requirement for a proline near the middle of the transmembrane domain of the murine leukemia virus (MLV) Env glycoprotein (Taylor and Sanders 1999).

The observations that mutations in fusion peptides or transmembrane domains (Armstrong et al. 2000; Baker et al. 1999; Tamm et al. 2002) can impair the ability to mediate full fusion (Fig. 3, Step 6) have suggested that both of these apolar domains function in the transition from hemifusion to full fusion. Initially, the fusion peptide appears to insert only into the outer leaflet of the target membrane (Tamm and Han 2000). It has been proposed that the transmembrane domain and the fusion peptide, which are close to each other after membrane merger, may interact to stabilize the fusion pore (Tamm et al. 2002; Zhou et al. 1997). If this is the case, the fusion peptide might span both leaflets of the fused membrane in its final conformation (Tamm et al. 2002).

6.3 The Juxtamembrane Region of the Ectodomain

Several lines of evidence suggest that ectodomain sequences that lie just before the transmembrane domains of certain viral fusion proteins may be important for fusion. These sequences tend to have a high proportion of tryptophans or other aromatic residues and are predicted to partition into the interfacial regions of membranes (Suarez et al. 2000). Indeed,
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synthetic peptides containing juxtamembrane ectodomain sequences from HIV Env and Ebola GP partition into the interfacial region of target membranes (Saez-Cirion et al. 2003; Saez-Cirion et al. 2002; Schibli et al. 2001; Suarez et al. 2000). Mutation of three tryptophans within this region of HIV gp41 abrogated infection (Salzwedel et al. 1999), apparently by inhibiting fusion pore enlargement (Munoz-Barroso et al. 1999). Extending the HIV gp41 C-terminal heptad repeat peptide to include the tryptophan-rich juxtamembrane ectodomain sequence appeared to increase the potency of the peptide as an inhibitor of fusion (Kliger et al. 2001). It was suggested that the extended heptad repeat peptide was more potent because it could bind to two sites on HIV Env (the N-terminal coiled-coil and a second, as yet unidentified site) (Kliger et al. 2001). A likely effect of these peptides on late stages of fusion is to prevent formation of a required structure in Env that provides additional membrane destabilization. In this manner, partitioning of juxtamembrane sequences into the interfacial region of membranes may promote the transition from a stalk intermediate to a fusion pore (see Sect. 5).

6.4 The Cytoplasmic Tail

A specific cytoplasmic tail sequence does not appear to be essential, but it can modulate late stages of fusion. The cytoplasmic tail has been shown to influence the transition from hemifusion to full fusion (Sakai et al. 2002) or fusion pore enlargement (Dutch and Lamb 2001; Kozerski et al. 2000) in some viruses. The mechanism by which cytoplasmic tails may influence these later stages of fusion is not known. Some studies using synthetic peptides have suggested a direct interaction between the cytoplasmic tail and the viral membrane (Chen et al. 2001b; Fujii et al. 1992; Gawrisch et al. 1993; Haffar et al. 1991; Kliger and Shai 1997). Others have shown that the cytoplasmic tail can influence the structure of the ectodomain of the fusion protein (Aguilar et al. 2003; Edwards et al. 2002).

The ability of the cytoplasmic tail to affect ectodomain structure is most clearly manifested for those viral fusion proteins that harbor fusion-suppressing sequences. These sequences have been found in the fusion proteins of MLV (Ragheb and Anderson 1994), other type C retroviruses (Bobkova et al. 2002), some lentiviruses (Kim et al. 2003; Luciw et al. 1998), and a paramyxovirus F protein (Tong et al. 2002). The cytoplasmic tail of MLV Env is cleaved during virus budding (Schultz and Rein 1985). MLV Envs harboring uncleaved cytoplasmic tails do not
induce fusion (Yang and Compans 1996). Although viruses lacking fusion-suppressing sequences display increased cell-cell fusion, they are more susceptible to neutralizing antibodies (Januszkesi et al. 1997; Li et al. 2001; Rein et al. 1994; Yang and Compans 1996) and are impaired in their ability to sustain multiple rounds of infection (Cathomen et al. 1998; Freed and Martin 1996; Piller et al. 2000).

Acylation of cytoplasmic tails can also affect fusion, apparently at a late stage. For example, a mutant HA from the Japan strain of influenza in which three (normally palmitoylated) cysteine residues were mutated appeared to fuse normally when monitored by dye redistribution assays (Melikyan et al. 1997b). However, electrophysiological measurements revealed that fusion pores formed by the mutant HA did not flicker like those formed by wt-HA (Melikyan et al. 1997b). Similar mutations in HA from the A/USSR/77 (H1N1) and A/FPV/Rostock/34 (H7N1) influenza subtypes were shown, respectively, to inhibit syncytia formation (Fischer et al. 1998) and the transition from hemifusion to full fusion (Sakai et al. 2002). Palmitoylation of HIV Env was also shown to be important for Env incorporation into virions and for infectivity (Rousso et al. 2000). Thus acylation of cytoplasmic tails appears to have multiple effects on viral fusion reactions, the details of which are not completely understood.

7
Rafts in Viral Membrane Fusion

Lipid rafts are plasma membrane microdomains that are enriched in cholesterol and glycosphingolipids with saturated acyl chains. They are organizational platforms for a variety of cellular functions including sorting of membrane proteins and signaling (Brown and London 2000). Although there is growing evidence that certain viruses employ rafts, or raftlike membrane microdomains, during virus assembly (Suomalainen 2002), the question of whether these structures are required at the site of fusion in the target cell is less clear. Here, we consider the role of rafts in the fusion of two enveloped viruses, SFV and HIV. It is important to consider whether cholesterol and/or sphingolipids are required for fusion because they are found in lipid rafts, or if they serve some other purpose. For example, cholesterol may interact directly with the fusion protein, thereby facilitating its insertion into the target membrane. Alternatively, a need for cholesterol and sphingolipids could reflect an ability of raft structures to concentrate viral receptors. A third possibility is that
the cholesterol imparts the membrane fluidity (or other biophysical properties) needed to lower the energy barrier for fusion.

SFV requires cholesterol and sphingolipids in the target membrane for fusion. These moieties enable the SFV spike protein to undergo conformational changes and bind to the target membrane (Ahn et al. 2002; Kielian et al. 2000). In a recent study, it was shown that after hydrophobic association with target bilayers, the SFV glycoprotein ectodomain associates with membrane structures with properties similar to rafts. However, careful studies using liposomes prepared with specific cholesterol and sphingolipid analogs demonstrated that the cholesterol and sphingolipid requirements in the target membrane did not correlate with their ability to form lipid rafts (Ahn et al. 2002). A related conclusion was drawn based on the fusion activities of both SFV and Sindbis virus with liposomes (Waarts et al. 2002). For both viruses, the requirement for cholesterol and sphingolipids in the target membrane appears to be for insertion of the fusion peptide (Vashishtha et al. 1998).

In the case of HIV, several studies have suggested a need for raft-like membrane microdomains for virus entry (Kozak et al. 2002; Popik et al. 2002). Depleting plasma membrane cholesterol from target cells resulted in reduced levels of virus infectivity or cell-cell fusion. Other studies have concluded that rafts are not necessary for HIV entry (Percherancier et al. 2003; Viard et al. 2002). In one study, depleting cholesterol from cells that express low levels of virus receptors inhibited HIV Env-mediated cell-cell fusion, but depleting cholesterol from cells that express high levels of virus receptors did not (Viard et al. 2002). Therefore, it was concluded that rafts per se are not needed for fusion. Rather, the presence of raft-like structures in the plasma membrane may concentrate virus receptors. Previous work has shown that a critical density of HIV receptors is required for fusion and infection (Reeves et al. 2002). Clearly more work is needed to clarify the role of rafts in virus-cell fusion and entry.

8 Inhibitors of Viral Fusion

It has recently become apparent, largely because of the success of T-20 in the inhibition of HIV infection in patients (Jiang et al. 2002; Kilby et al. 1998), that fusion is a good target for antiviral intervention. This was originally conceptualized because fusion is an essential early step in the virus infectious cycle, it happens in an exoplasmic space, and strategies
can be designed to inhibit fusion without interfering with host cell proteins. Some fusion inhibitors function by inhibiting six-helix bundle formation. Others function by preventing earlier conformational changes in viral fusion proteins.

### 8.1 Inhibition of Helix Bundle Formation

The peptide T-20 (also known as Fuzeon) corresponds to the C-terminal helix of HIV Env (Fig. 3, green). T-20 works by preventing six-helix bundle formation. T-20 is a potent inhibitor of infections in tissue culture. Peptides corresponding to equivalent regions of other retroviruses as well as several paramyxoviruses function similarly (Earp et al. 2003; Russell et al. 2001). Notably, all of the viruses that have been shown to be highly susceptible to “C-helix” peptide inhibitors function at neutral pH, at least up to the lipid interacting stage of virus-cell fusion (Earp et al. 2003). Peptides corresponding to the N-terminal helices of HIV Env and the SV5 F protein also inhibit fusion, although with lower potency (Lu et al. 1995; Russell et al. 2001). The mechanism of inhibition by N-terminal peptides is still under consideration (He et al. 2003). In the case of the SV5 F, the N-peptide appears to target an earlier intermediate than the C-peptide (Russell et al. 2001). Other strategies are being considered to stabilize the prehairpin intermediate (Fig. 3, Step 1) and thereby prevent six-helix bundle formation. One strategy is the development of antibodies that recognize the prehairpin intermediate (Golding et al. 2002). Another, exemplified in three studies, is the development of small molecules that prevent six-helix bundle formation (Debnath et al. 1999; Eckert et al. 1999; Ferrer et al. 1999). All three studies targeted a hydrophobic pocket in the groove of the central coiled-coil of HIV gp41 that is important for interaction with the C-terminal helix in the post-fusion form. In the first approach, two organic compounds were identified from a screening effort conducted in conjunction with molecular docking, a method to identify small molecules that fit in a target site (Debnath et al. 1999). The second approach replaced three residues of the C-terminal helix that bind to the hydrophobic pocket with organic moieties, generated by combinatorial chemistry (Ferrer et al. 1999). The third approach used a mirror image phage display library to identify small, D-amino acid containing peptides that bind to the pocket (Eckert et al. 1999). Although none of the small molecules identified to date is as potent as T-20, the precedent has been set for attaining this goal.
8.2 Inhibition of Other Steps in Fusion

An effort to block the fusion activity of influenza was based on the idea that maintaining HA in its native metastable state should prevent fusion and infection. The first trial targeted a site in X:31 HA that includes part of the fusion peptide. With the use of an antibody-based assay to monitor fusion peptide exposure, a compound, tert-butylhydroquinone (TBHQ), that prevents the first stages of the HA conformational change and inhibits infectivity was discovered (Bodian et al. 1993). A follow-up study, targeting a site near the B-loop in HA, yielded additional inhibitors. Whereas some functioned like TBHQ, a second class was identified that appeared to push HA to an inactive state (Hoffman et al. 1997). A random screen against an H1 influenza virus identified an inhibitor that appears to function similarly to TBHQ. In the latter case, the binding site for the inhibitor was mapped to the vicinity of the fusion peptide (Cianci et al. 1999). Other small molecules that inhibit conformational changes in HA have been identified (Staschke et al. 1998). To date, none of the HA inhibitors has blocked all HA subtypes and none has an IC$_{50}$ value in the submicromolar range. It is not yet clear whether the latter limitation represents a fundamental difficulty in inhibiting viral fusion proteins that function at low pH.

In addition to the small molecule approaches described above, antibodies that prevent fusion-inducing changes in viral glycoproteins have been described. The first example was an antibody that prevents low-pH-induced fusion of West Nile virus with model liposomes (Gollins and Porterfield 1986). Recently, a Fab fragment that binds to two HA1 monomers was shown to prevent an early conformational change in influenza HA (Barbey-Martin et al. 2002), separation of the globular head domains (Fig. 2B). As described above, antibodies have been developed that likely block six-helix bundle formation in the case of HIV Env (Golding et al. 2002).

9 Perspectives

The goal of this review was to give the reader an appreciation for the diversity of viral fusion mechanisms while recognizing their common underlying principles. We also summarized what is known about the lipid dynamics and lipid structures involved in fusion, and we also briefly
overviewed recent developments in targeting viral fusion as an antiviral strategy. There is clearly much more we need to know about viral fusion proteins, viral fusion reactions, and the design of antifusion agents.

We end this review by enumerating some pressing issues and questions that remain about viral fusion. A major goal is to determine high-resolution structures for the complete ectodomains of the metastable trimers of class I viral fusion proteins in addition to the influenza HA. Structures of a complete paramyxovirus F and a complete retroviral Env ectodomain will be highly informative because we currently lack a detailed molecular description of how a receptor activates any viral fusion protein at neutral pH. A second goal will be to further delineate the mechanisms of class II viral fusion proteins. Do the transitions to their recently described low pH forms (Bressanelli et al. 2004; Gibbons et al. 2004; Modis et al. 2004) mediate hemifusion or fusion pore opening? What about the mechanisms of the as yet unclassified viral fusion proteins? These include viruses such as rhabdoviruses (e.g., VSV) that need only one protein to promote fusion, as well as more complicated viruses such as herpesviruses and poxviruses that require multiple viral glycoproteins.

The ensuing years should also bring a more complete understanding of how viral fusion proteins interact with target membrane bilayers. Class II fusion proteins insert their internal fusion peptides into target membranes as loops (Bressanelli et al. 2004; Gibbons et al. 2004; Modis et al. 2004). It has been predicted that the internal fusion proteins of the class I fusion proteins from Ebola and avian retroviruses form disulfide-bonded loop structures (Weisenhorn et al. 1998), and mutagenesis work has supported this prediction (Delos et al. 2000; Delos and White 2000; Jeffers et al. 2002). It remains to be seen, from high resolution structural studies, whether all internal fusion peptides, be they from class I, class II, or other classes of fusion proteins, interact with target bilayers as (disulfide bond) stabilized loops. Finally, we expect that there will be major developments in furthering the concept of targeting fusion as a weapon against pathogenic enveloped viruses. Particular emphasis will likely be on the development of small molecule inhibitors through the use of combinatorial chemistry in conjunction with high-throughput screens. It will be interesting to learn whether small molecule fusion inhibitors can be identified that block the entry of viruses that fuse in endosomes in response to low pH. This is a challenge for low-pH-activated class I fusion proteins such as influenza HA as well as for all known class II fusion proteins. Stay tuned. There are likely to be exciting develop-
ments in our understanding of viral fusion mechanisms as well as in the development of antifusion antivirals in the years ahead.

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**References**

Abrahamyan LG, Markosyan RM, Moore JP, Cohen FS, and Melikyan GB (2003) Human immunodeficiency virus type 1 Env with an intersubunit disulfide bond engages coreceptors but requires bond reduction after engagement to induce fusion. J Virol 77:5829–36

Aguilar HC, Anderson WF, and Cannon PM (2003) Cytoplasmic tail of moloney murine leukemia virus envelope protein influences the conformation of the extracellular domain: implications for the mechanism of action of the R peptide. J Virol 77:1281–1291

Ahn A, Gibbons DL, and Kielian M (2002) The fusion peptide of Semliki Forest virus associates with sterol-rich membrane domains. J Virol 76:10029–39

Ahn A, Klimjack MR, Chatterjee PK, and Kielian M (1999) An epitope of the Semliki Forest virus fusion protein exposed during virus-membrane fusion. J Virol 73:5605–12

Allison SL, Schalich J, Stiasny K, Mandl CW, Kunz C, and Heinz FX (1995) Oligomeric rearrangement of tick-borne encephalitis virus envelope proteins induced by an acidic pH. J Virol 69:695–700

Allison SL, Stiasny K, Stadler K, Mandl CW, and Heinz FX (1999) Mapping of functional elements in the stem-anchor region of tick-borne encephalitis virus envelope protein E. J Virol 73:5605–12

Allison SL, Schalich J, Stiasny K, Mandl CW, and Heinz FX (2001) Mutational evidence for an internal fusion peptide in flavivirus envelope protein E. J Virol 75:4268–75

Armstrong RT, Kushnir AS, and White JM (2000) The transmembrane domain of influenza hemagglutinin exhibits a stringent length requirement to support the hemifusion to fusion transition. J Cell Biol 151:425–38

Bagai S and Lamb RA (1995) Quantitative measurement of paramyxovirus fusion: differences in requirements of glycoproteins between simian virus 5 and human parainfluenza virus 3 or Newcastle disease virus. J Virol 69:6712–9

Baker KA, Dutch RE, Lamb RA, and Jardetzky TS (1999) Structural basis for paramyxovirus-mediated membrane fusion. Mol Cell 3:309–19

Barbey-Martín C, Gigant B, Bizebard T, Calder LJ, Wharton SA, Skehel JJ, and Knowsow M (2002) An antibody that prevents the hemagglutinin low pH fusogenic transition. Virology 294:70–4

Barbouche R, Miqelis R, Jones IM, and Fenouillet E (2003) Protein-disulfide isomerase-mediated reduction of two disulfide bonds of HIV envelope glycoprotein 120 occurs post-CXCR4 binding and is required for fusion. J Biol Chem 278:3131–3136
Barnett AL, and Cunningham JM (2001) Receptor binding transforms the surface subunit of the mammalian C-type retrovirus envelope protein from an inhibitor to an activator of fusion. J Virol 75:9096–105

Begona Ruiz-Arguello M, Gonzalez-Reyes L, Calder LJ, Palomo C, Martin D, Saiz MJ, Garcia-Barreno B, Skehel JJ, and Melero JA (2002) Effect of proteolytic processing at two distinct sites on shape and aggregation of an anchorless fusion protein of human respiratory syncytial virus and fate of the intervening segment. Virology 298:317–26

Blumenthal R, Sarkar DP, Durell S, Howard DE, and Morris SJ (1996) Dilation of the influenza hemagglutinin fusion pore revealed by the kinetics of individual cell-fusion events. J Cell Biol 135:63–71

Bobkova M, Stitz J, Engelstädter M, Cichutek K, and Buchholz CJ (2002) Identification of R-peptides in envelope proteins of C-type retroviruses. J Gen Virol 83:2241–6

Bodian DL, Yamasaki RB, Buswell RL, Stearns JF, White JM, and Kuntz ID (1993) Inhibition of the fusion-inducing conformational change of influenza hemagglutinin by benzoquinones and hydroquinones. Biochemistry 32:2967–78

Borrego-Diaz E, Peeples ME, Marksosyan RM, Melikyan GB, and Cohen FS (2003) Completion of trimeric hairpin formation of influenza virus hemagglutinin promotes fusion pore opening and enlargement. Virology 316:234–44

Bosch BJ, Zee R van der, Haan CA de, and Rottier PJ (2003) The coronavirus spike protein is a class I virus fusion protein: structural and functional characterization of the fusion core complex. J Virol 77:8801–11

Bossart KN, Wang LF, Flora MN, Chua KB, Lam SK, Eaton BT, and Broder CC (2002) Membrane fusion tropism and heterotypic functional activities of the nipa virus and hendra virus envelope glycoproteins. J Virol 76:11186–11198

Bressanelli S, Stiasny K, Allison SL, Stura EA, Duquerroy S, Lescar J, Heinz FX, and Ray FA (2004) Structure of a flavivirus envelope glycoprotein in its low-pH-induced membrane fusion conformation. EMBO J 23:728–38

Bron R, Wahlberg JM, Garoff H, and Wilschut J (1993) Membrane fusion of Semliki Forest virus in a model system: correlation between fusion kinetics and structural changes in the envelope glycoprotein. EMBO J 12:693–701

Brown DA and London E (2000) Structure and function of sphingolipid- and cholesterol-rich membrane rafts. J Biol Chem 275:17221–4

Bullough PA, Hughson FM, Skehel JJ, and Wiley DC (1994) Structure of influenza haemagglutinin at the pH of membrane fusion. Nature 371:37–43

Calder LJ, Gonzalez-Reyes L, Garcia-Barreno B, Wharton SA, Skehel JJ, Wiley DC, and Melero JA (2000) Electron microscopy of the human respiratory syncytial virus fusion protein and complexes that it forms with monoclonal antibodies. Virology 271:122–31

Carr CM, Chaudhry C, and Kim PS (1997) Influenza hemagglutinin is spring-loaded by a metastable native conformation. Proc Natl Acad Sci USA 94:14306–13

Carr CM and Kim PS (1993) A spring-loaded mechanism for the conformational change of influenza hemagglutinin. Cell 73:823–32

Cathomen T, Naim HY, and Cattaneo R (1998) Measles viruses with altered envelope protein cytoplasmic tails gain cell fusion competence. J Virol 72:1224–34
Chan DC, Fass D, Berger JM, and Kim PS (1997) Core structure of gp41 from the HIV envelope glycoprotein. Cell 89:263–73
Chan DC and Kim PS (1998) HIV entry and its inhibition. Cell 93:681–4
Chen J, Lee KH, Steinhauer DA, Stevens DJ, Skehel JJ, and Wiley DC (1998) Structure of the hemagglutinin precursor cleavage site, a determinant of influenza pathogenicity and the origin of the labile conformation. Cell 95:409–17
Chen J, Skehel JJ, and Wiley DC (1999) N- and C-terminal residues combine in the fusion-pH influenza hemagglutinin HA(2) subunit to form an N cap that terminates the triple-stranded coiled coil. Proc Natl Acad Sci USA 96:8967–72
Chen L, Gorman JJ, McKimm-Breschkin J, Lawrence LJ, Tulloch PA, Smith BJ, Colman PM, and Lawrence MC (2001a) The structure of the fusion glycoprotein of Newcastle disease virus suggests a novel paradigm for the molecular mechanism of membrane fusion. Structure 9:255–66
Chen SS, Lee SF, and Wang CT (2001b) Cellular membrane-binding ability of the C-terminal cytoplasmic domain of human immunodeficiency virus type 1 envelope transmembrane protein gp41. J Virol 75:9925–38
Cianci C, Yu KL, Dischino DD, Harte W, Deshpande M, Luo G, Colomno RJ, Meanwell NA, and Krystal M (1999) pH-dependent changes in photoaffinity labeling patterns of the H1 influenza virus hemagglutinin by using an inhibitor of viral fusion. J Virol 73:1785–94
Cleverley DZ and Lenard J (1998) The transmembrane domain in viral fusion: essential role for a conserved glycine residue in vesicular stomatitis virus G protein. Proc Natl Acad Sci USA 95:3425–30
Colman PM and Lawrence MC (2003) The structural biology of type I viral membrane fusion. Nat Rev Mol Cell Biol 4:309–19
Corver J, Ortiz A, Allison SL, Schalich J, Heinz FX, and Wilschut J (2000) Membrane fusion activity of tick-borne encephalitis virus and recombinant subviral particles in a liposomal model system. Virology 269:37–46
Damico RL, Crane J, and Bates P (1998) Receptor-triggered membrane association of a model retroviral glycoprotein. Proc Natl Acad Sci USA 95:2580–5
Danieli T, Pelletier SL, Henis YI, and White JM (1996) Membrane fusion mediated by the influenza virus hemagglutinin requires the concerted action of at least three hemagglutinin trimers. J Cell Biol 133:559–69
Debnath AK, Radigan L, and Jiang S (1999) Structure-based identification of small molecule antiviral compounds targeted to the gp41 core structure of the human immunodeficiency virus type 1. J Med Chem 42:3203–9
Delos SE, Gilbert JM, and White JM (2000) The central proline of an internal viral fusion peptide serves two important roles. J Virol 74:1686–93
Delos SE and White JM (2000) Critical role for the cysteines flanking the internal fusion peptide of avian sarcoma/leukosis virus envelope glycoprotein. J Virol 74:9738–41
Deng R, Wang Z, Mahon PJ, Marinello M, Mirza A, and Iorio RM (1999) Mutations in the Newcastle disease virus hemagglutinin-neuraminidase protein that interfere with its ability to interact with the homologous F protein in the promotion of fusion. Virology 253:43–54
Dennison SM, Greenfield N, Lenard J, and Lentz BR (2002) VSV transmembrane domain (TMD) peptide promotes PEG-mediated fusion of liposomes in a conformationally sensitive fashion. Biochemistry 41:14925–14934

Di Simone C and Buchmeier MJ (1995) Kinetics and pH dependence of acid-induced structural changes in the lymphocytic choriomeningitis virus glycoprotein complex. Virology 209:3–9

Doms RW, Helenius A, and White J (1985) Membrane fusion activity of the influenza virus hemagglutinin. The low pH-induced conformational change. J Biol Chem 260:2973–81

Duffus WA, Levy-Mintz P, Klimjack MR, and Kielian M (1995) Mutations in the putative fusion peptide of Semliki Forest virus affect spike protein oligomerization and virus assembly. J Virol 69:2471–9

Durell SR, Martin I, Ruysschaert JM, Shai Y, and Blumenthal R (1997) What studies of fusion peptides tell us about viral envelope glycoprotein-mediated membrane fusion. Mol Membr Biol 14:97–112

Dutch RE and Lamb RA (2001) Deletion of the cytoplasmic tail of the fusion protein of the paramyxovirus simian virus 5 affects fusion pore enlargement. J Virol 75:5363–9

Earp LJ, Delos SE, Netter RC, Bates P, and White JM (2003) The avian retrovirus avian sarcoma/leukosis virus subtype A reaches the lipid mixing stage of fusion at neutral pH. J Virol 77:3058–3066

Eckert DM and Kim PS (2001) Mechanisms of viral membrane fusion and its inhibition. Annu Rev Biochem 70:777–810

Eckert DM, Malashkevich VN, Hong LH, Carr PA, and Kim PS (1999) Inhibiting HIV-1 entry: discovery of D-peptide inhibitors that target the gp41 coiled-coil pocket. Cell 99:103–15

Edwards TG, Wyss S, Reeves JD, Zolla-Pazner S, Hoxie JA, Doms RW, and Baribaud F (2002) Truncation of the cytoplasmic domain induces exposure of conserved regions in the ectodomain of human immunodeficiency virus type 1 envelope protein. J Virol 76:2683–91

Einfeld DA and Hunter E (1994) Expression of the TM protein of Rous sarcoma virus in the absence of SU shows that this domain is capable of oligomerization and intracellular transport. J Virol 68:2513–20

Fackler OT and Peterlin BM (2000) Endocytic entry of HIV-1. Curr Biol 10:1005–8

Feng Y, Broder CC, Kennedy PE, and Berger EA (1996) HIV-1 entry cofactor: functional cDNA cloning of a seven-transmembrane, G protein-coupled receptor. Science 272:872–7

Fenouillet E, Barbouche R, Courageot J, and Miquelis R (2001) The catalytic activity of protein disulfide isomerase is involved in human immunodeficiency virus envelope-mediated membrane fusion after CD4 cell binding. J Infect Dis 183:744–52

Ferlenghi I, Clarke M, Ruttan T, Allison SL, Schalich J, Heinz FX, Harrison SC, Rey F, and Fuller SD (2001) Molecular organization of a recombinant subviral particle from tick-borne encephalitis virus. Mol Cell 7:593–602

Ferrr M, Kapoor TM, Strassmaier T, Weissenhorn W, Skehel JJ, Oprian D, Schreiber SL, Wiley DC, and Harrison SC (1999) Selection of gp41-mediated HIV-1 cell en-
try inhibitors from biased combinatorial libraries of non-natural binding elements. Nat Struct Biol 6:953–60
Fischer C, Schroth-Diez B, Herrmann A, Garten W, and Klenk HD (1998) Acylation of the influenza hemagglutinin modulates fusion activity. Virology 248:284–94
Fredericksen BL and Whitt MA (1995) Vesicular stomatitis virus glycoprotein mutations that affect membrane fusion activity and abolish virus infectivity. J Virol 69:1435–43
Freed EO, Delwart EL, Buchschacher GLJ, and Panganiban AT (1992) A mutation in the human immunodeficiency virus type 1 transmembrane glycoprotein gp41 dominantly interferes with fusion and infectivity. Proc Natl Acad Sci USA 89:70–4
Freed EO and Martin MA (1996) Domains of the human immunodeficiency virus type 1 matrix and gp41 cytoplasmic tail required for envelope incorporation into virions. J Virol 70:341–51
Frey S, Marsh M, Gunther S, Pelchen-Matthews A, Stephens P, Ortlepp S, and Stegmann T (1995) Temperature dependence of cell-cell fusion induced by the envelope glycoprotein of human immunodeficiency virus type 1. J Virol 69:1462–72
Fujii G, Horvath S, Woodward S, Eiserling F, and Eisenberg D (1992) A molecular model for membrane fusion based on solution studies of an amphiphilic peptide from HIV gp41. Protein Sci 1:1454–64
Furuta RA, Wild CT, Weng Y, and Weiss CD (1998) Capture of an early fusion-active conformation of HIV-1 gp41. Nat Struct Biol 5:276–9
Gallina A, Hanley TM, Mandel R, Trahey M, Broder CC, Viglianti GA, and Ryser HJ (2002) Inhibitors of protein-disulfide isomerase prevent cleavage of disulfide bonds in receptor-bound glycoprotein 120 and prevent HIV-1 entry. J Biol Chem 277:50579–88
Gaudin Y, de Kinkelin P, and Benmansour A (1999a) Mutations in the glycoprotein of viral haemorrhagic septicemia virus that affect virulence for fish and the pH threshold for membrane fusion. J Gen Virol 80:1221–9
Gaudin Y, Tuffereau C, Durrer P, Brunner J, Flamand A, and Ruigrok R (1999b) Rabies virus-induced membrane fusion. Mol Membr Biol 16:21–31
Gawrisch K, Han KH, Yang JS, Bergelson LD, and Ferretti JA (1993) Interaction of peptide fragment 828–848 of the envelope glycoprotein of human immunodeficiency virus type I with lipid bilayers. Biochemistry 32:3112–8
Gething MJ, Doms RW, York D, and White J (1986) Studies on the mechanism of membrane fusion: site-specific mutagenesis of the hemagglutinin of influenza virus. J Cell Biol 102:11–23
Gibbons DL, Vaney MC, Roussel A, Vigouroux A, Reilly B, Lepault J, Kielian M, and Rey FA (2004) Conformational change and protein-protein interactions of the fusion protein of Semliki Forest virus. Nature 427:320–25
Gilbert JM, Mason D, and White JM (1990) Fusion of Rous sarcoma virus with host cells does not require exposure to low pH. J Virol 64:5106–13
Godley L, Pfeifer J, Steinhauer D, Ely B, Shaw G, Kaufmann R, Suchanek E, Pabo C, Skehel JJ, and Wiley DC (1992) Introduction of intersubunit disulfide bonds in the membrane-distal region of the influenza hemagglutinin abolishes membrane fusion activity. Cell 68:635–45
Golding H, Zaitseva M, de Rosny E, King LR, Manischewitz J, Sidorov I, Gorny MK, Zolla-Pazner S, Dimitrov DS, and Weiss C (2002) Dissection of human immunodeficiency virus type 1 entry with neutralizing antibodies to gp41 fusion intermediates. J Virol 76:6780–90

Gollins SW and Porterfield JS (1986) A new mechanism for the neutralization of enveloped viruses by antiviral antibody. Nature 321:244–6

Gonzalez-Reyes L, Ruiz-Arguello MB, Garcia-Barreno B, Calder L, Lopez JA, Albar JP, Skehel JJ, Wiley DC, and Melero JA (2001) Cleavage of the human respiratory syncytial virus fusion protein at two distinct sites is required for activation of membrane fusion. Proc Natl Acad Sci USA 98:9859–64

Gruenke JA, Armstrong RT, Newcomb WW, Brown JC, and White JM (2002) New insights into the spring-loaded conformational change of influenza hemagglutinin. J Virol 76:4456–66

Gunther-Ausborn S, Schoen P, Bartoldus I, Wilschut J, and Stegmann T (2000) Role of hemagglutinin surface density in the initial stages of influenza virus fusion: lack of evidence for cooperativity. J Virol 74:2714–20

Haffar OK, Dowbenko DJ, and Berman PW (1991) The cytoplasmic tail of HIV-1 gp160 contains regions that associate with cellular membranes. Virology 180:439–41

Han X, Bushweller JH, Cafiso DS, and Tamm LK (2001) Membrane structure and fusion-triggering conformational change of the fusion domain from influenza hemagglutinin. Nat Struct Biol 8:715–20

He Y, Vassell R, Zaitseva M, Nguyen N, Yang Z, Weng Y, and Weiss C (2003) Peptides trap the human immunodeficiency virus type 1 envelope glycoprotein fusion intermediate at two sites. J Virol 77:1666–1671

Heinz FX and Allison SL (2001) The machinery for flavivirus fusion with host cell membranes. Curr Opin Microbiol 4:450–5

Helenius A (1995) Alphavirus and flavivirus glycoproteins: structures and functions. Cell 81:651–3

Hernandez LD, Hoffman LR, Wolfsberg TG, and White JM (1996) Virus-cell and cell-cell fusion. Annu Rev Cell Dev Biol 12:627–61

Hernandez LD, Peters RJ, Delos SE, Young JA, Agard DA, and White JM (1997) Activation of a retroviral membrane fusion protein: soluble receptor-induced liposome binding of the ALSV envelope glycoprotein. J Cell Biol 139:1455–54

Hernandez LD and White JM (1998) Mutational analysis of the candidate internal fusion peptide of the avian leukosis and sarcoma virus subgroup A envelope glycoprotein. J Virol 72:3259–67

Hoffman LR, Kuntz ID, and White JM (1997) Structure-based identification of an inducer of the low-pH conformational change in the influenza virus hemagglutinin: irreversible inhibition of infectivity. J Virol 71:8808–20

Hunter E (1997) Viral entry and receptors. In J. M. Coffin (ed.), Retroviruses. Cold Spring Harbor Laboratory Press, Plainview, NY, p. 71–121

Irurzun A, Nieva JL, and Carrasco L (1997) Entry of Semliki forest virus into cells: effects of concanamycin A and nigericin on viral membrane fusion and infection. Virology 227:488–92

Ito H, Watanabe S, Sanchez A, Whitt MA, and Kawaoka Y (1999) Mutational analysis of the putative fusion domain of Ebola virus glycoprotein. J Virol 73:8907–12
Jahn R, Lang T, and Sudhof TC (2003) Membrane fusion. Cell 112:519–533
Januszkeski MM, Cannon PM, Chen D, Rozenberg Y, and Anderson WF (1997) Functional analysis of the cytoplasmic tail of Moloney murine leukemia virus envelope protein. J Virol 71:3613–9
Jeffers SA, Sanders DA, and Sanchez A (2002) Covalent modifications of the ebola virus glycoprotein. J Virol 76:12463–72
Jiang S, Zhao Q, and Debnath AK (2002) Peptide and non-peptide HIV fusion inhibitors. Curr Pharm Des 8:563–80
Joshi SB, Dutch RE, and Lamb RA (1998) A core trimer of the paramyxovirus fusion protein: parallels to influenza virus hemagglutinin and HIV-1 gp41. Virology 248:20–34
Kamath S and Wong TC (2002) Membrane structure of the human immunodeficiency virus gp41 fusion domain by molecular dynamics simulation. Biophys J 83:135–43
Kanaseki T, Kawasaki K, Murata M, Ikeuchi Y, and Ohnishi S (1997) Structural features of membrane fusion between influenza virus and liposome as revealed by quick-freezing electron microscopy. J Cell Biol 137:1041–56
Kemble GW, Bodian DL, Rose J, Wilson IA, and White JM (1992) Intermonomer disulfide bonds impair the fusion activity of influenza virus hemagglutinin. J Virol 66:4940–50
Kemble GW, Danieli T, and White JM (1994) Lipid-anchored influenza hemagglutinin promotes hemifusion, not complete fusion. Cell 76:383–91
Kielian M (1995) Membrane fusion and the alphavirus life cycle. Adv Virus Res 45:113–51
Kielian M, Klimjack MR, Ghosh S, and Duffus WA (1996) Mechanisms of mutations inhibiting fusion and infection by Semliki Forest virus. J Cell Biol 134:863–72
Kielian M, Chatterjee PK, Gibbons DL, and Lu YE (2000) Specific roles for lipids in virus fusion and exit. Examples from the alphaviruses. Subcell Biochem 34:409–55
Kilby JM, Hopkins S, Venetta TM, DDiMassimo B, Cloud GA, Lee JY, Alldredge L, Hunter E, Lambert D, Bolognesi D, Matthews T, Johnson MR, Nowak MA, Shaw GM, and Saag MS (1998) Potent suppression of HIV-1 replication in humans by T-20, a peptide inhibitor of gp41-mediated virus entry. Nat Med 4:1302–7
Kim FJ, Manel N, Boublik Y, Battini JL, and Sitbon M (2003) Human T-cell leukemia virus type 1 envelope-mediated syncytium formation can be activated in resistant mammalian cell lines by a carboxy-terminal truncation of the envelope cytoplasmic domain. J Virol 77:963–969
Kliger Y, Aharoni A, Rapaport D, Jones P, Blumenthal R, and Shai Y (1997) Fusion peptides derived from the HIV type 1 glycoprotein 41 associate within phospholipid membranes and inhibit cell-cell fusion. Structure-function study. J Biol Chem 272:13496–505
Kliger Y, Gallo SA, Peisajovich SG, Munoz-Barroso I, Avkin S, Blumenthal R, and Shai Y (2001) Mode of action of an antiviral peptide from HIV-1. Inhibition at a post-lipid mixing stage. J Biol Chem 276:1391–7
Kliger Y and Shai Y (1997) A leucine zipper-like sequence from the cytoplasmic tail of the HIV-1 envelope glycoprotein binds and perturbs lipid bilayers. Biochemistry 36:5157–69
Korte T, Ludwig K, Booy FP, Blumenthal R, and Herrmann A (1999) Conformational intermediates and fusion activity of influenza virus hemagglutinin. J Virol 73:4567–74
Kozak SL, Heard JM, and Kabat D (2002) Segregation of CD4 and CXCR4 into distinct lipid microdomains in T lymphocytes suggests a mechanism for membrane destabilization by human immunodeficiency virus. J Virol 76:1802–15
Kozerski C, Ponimaskin E, Schroth-Diez B, Schmidt MF, and Herrmann A (2000) Modification of the cytoplasmic domain of influenza virus hemagglutinin affects enlargement of the fusion pore. J Virol 74:7529–37
Kozlov MM and Chernomordik LV (2002) The protein coat in membrane fusion: lessons from fission. Traffic 3:256–67
Kozlovsky Y and Kozlov MM (2002) Stalk model of membrane fusion: solution of energy crisis. Biophys J 82:882–95
Kuhn RJ, Zhang W, Rossmann MG, Pletnev SV, Corver J, Lenches E, Jones CT, Mukhopadhyay S, Chipman PR, Strauss EG, Baker TS, and Strauss JH (2002) Structure of dengue virus: implications for flavivirus organization, maturation, and fusion. Cell 108:717–25
Kwong PD, Wyatt R, Robinson J, Sweet RW, Sodroski J, and Hendrickson W (1998) Structure of an HIV gp120 envelope glycoprotein in complex with the CD4 receptor and a neutralizing human antibody. Nature 393:648–59
Lamb RA (1993) Paramyxovirus fusion: a hypothesis for changes. Virology 197:1-11
Lambert DM, Barney S, Lambert AL, Guthrie K, Medina R, Davis DE, Bucy T, Erickson J, Merutka G, and Petteway SRJ (1996) Peptides from conserved regions of paramyxovirus fusion (F) proteins are potent inhibitors of viral fusion. Proc Natl Acad Sci USA 93:2186–91
Lavillette D, Boson B, Russell SJ, and Cosset FL (2001) Activation of membrane fusion by murine leukemia viruses is controlled in cis or in trans by interactions between the receptor-binding domain and a conserved disulfide loop of the carboxy terminus of the surface glycoprotein. J Virol 75:3685–95
Lavillette D, Maurice M, Roche C, Russell SJ, Sitbon M, and Cosset FL (1998) A proline-rich motif downstream of the receptor binding domain modulates conformation and fusogenicity of murine retroviral envelopes. J Virol 72:9955–65
Lescar J, Roussel A, Wien MW, Navaza J, Fuller SD, Wengler G, and Rey FA (2001) The fusion glycoprotein shell of Semliki Forest virus: an icosahedral assembly primed for fusogenic activation at endosomal pH. Cell 105:137–48
Li M, Yang C, and Compans RW (2001) Mutations in the cytoplasmic tail of murine leukemia virus envelope protein suppress fusion inhibition by R peptide. J Virol 75:2337–44
Li Y, Drone C, Sat E, and Ghosh HP (1993) Mutation analysis of the vesicular stomatitis virus glycoprotein G for membrane fusion domains. J Virol 67:4070–7
Li Y, Han X, and Tamm NK (2003) Thermodynamics of fusion peptide-membrane interactions. Biochemistry 42:27245–51
Lu M, Blacklow SC, and Kim PS (1995) A trimeric structural domain of the HIV-1 transmembrane glycoprotein. Nat Struct Biol 2:1075–82
Luciw PA, Shaw KE, Shacklett BL, and Marthas ML (1998) Importance of the intracytoplasmic domain of the simian immunodeficiency virus (SIV) envelope glycoprotein for pathogenesis. Virology 252:9–16
Markosyan RM, Cohen FS, and Melikyan GB (2003) HIV-1 envelope proteins complete their folding into six-helix bundles immediately after fusion pore formation. Mol Biol Cell 14:926–38

Markovic I, Leikina E, Zhukovsky M, Zimmerberg J, and Chernomordik LV (2001) Synchronized activation and refolding of influenza hemagglutinin in multimeric fusion machines. J Cell Biol 155:833–44

Markovic I, Pulyaeva H, Sokoloff A, and Chernomordik LV (1998) Membrane fusion mediated by baculovirus gp64 involves assembly of stable gp64 trimers into multiprotein aggregates. J Cell Biol 143:1155–66

Martin I and Ruysschaert JM (2000) Common properties of fusion peptides from diverse systems. Biosci Rep 20:483–500

Martin II, Ruysschaert J, and Epand RM (1999) Role of the N-terminal peptides of viral envelope proteins in membrane fusion. Adv Drug Deliv Rev 38:233–255

Matsuyama S, Delos SE, and White JM (2004) Sequential roles of receptor binding and low pH in forming prehairpin and hairpin conformations of an avian retroviral envelope glycoprotein. J Virol (in press)

McClure MO, Sommerfelt MA, Marsh M, and Weiss RA (1990) The pH independence of mammalian retrovirus infection. J Gen Virol 71:767–73

McGinnes LW, Gravel K, and Morrison TG (2002) Newcastle disease virus HN protein alters the conformation of the F protein at cell surfaces. J Virol 76:12622–33

Melikyan GB, Brener SA, Ok DC, and Cohen FS (1997a) Inner but not outer membrane leaflets control the transition from glycosylphosphatidylinositol-anchored influenza hemagglutinin-induced hemifusion to full fusion. J Cell Biol 136:995–1005

Melikyan GB, Jin H, Lamb RA, and Cohen FS (1997b) The role of the cytoplasmic tail region of influenza virus hemagglutinin in formation and growth of fusion pores. Virology 235:118–28

Melikyan GB, Markosyan RM, Hemmati H, Delmedico MK, Lambert DM, and Cohen FS (2000a) Evidence that the transition of HIV-1 gp41 into a six-helix bundle, not the bundle configuration, induces membrane fusion. J Cell Biol 151:413–23

Melikyan GB, Markosyan RM, Roth MG, and Cohen FS (2000b) A point mutation in the transmembrane domain of the hemagglutinin of influenza virus stabilizes a hemifusion intermediate that can transit to fusion. Mol Biol Cell 11:3765–75

Modis Y, Ogata S, Clements D, and Harrison SC (2004) Structure of the dengue virus envelope protein after membrane fusion. Nature 427:313–9

Mothes W, Boerger AL, Narayan S, Cunningham JM, and Young JA (2000) Retroviral entry mediated by receptor priming and low pH triggering of an envelope glycoprotein. Cell 103:679–89

Munoz-Barroso I, Durrell S, Sakaguchi K, Appella E, and Blumenthal R (1998) Dilation of the human immunodeficiency virus-1 envelope glycoprotein fusion pore revealed by the inhibitory action of a synthetic peptide from gp41. J Cell Biol 140:315–23

Munoz-Barroso I, Salzwedel K, Hunter E, and Blumenthal R (1999) Role of the membrane-proximal domain in the initial stages of human immunodeficiency virus type 1 envelope glycoprotein-mediated membrane fusion. J Virol 73:6089–92

Netter RC (2002) Ph.D Dissertation. University of Pennsylvania.
Nir S, Duzgunes N, de Lima MC, and Hoekstra D (1990) Fusion of enveloped viruses with cells and liposomes. Activity and inactivation. Cell Biophys 17:181–201
Nussbaum O, Broder CC, and Berger EA (1994) Fusogenic mechanisms of enveloped-virus glycoproteins analyzed by a novel recombinant vaccinia virus-based assay quantitating cell fusion-dependent reporter gene activation. J Virol 68:5411–22
Ohuchi M, Ohuchi R, Sakai T, and Matsumoto A (2002) Tight binding of influenza virus hemagglutinin to its receptor interferes with fusion pore dilation. J Virol 76:1405-13
Owens RJ, Burke C, and Rose JK (1994) Mutations in the membrane-spanning domain of the human immunodeficiency virus envelope glycoprotein that affect fusion activity. J Virol 68:570–4
Park HE, Gruenke JA, and White JM (2003) Leash in the groove mechanism of membrane fusion. Nat Struct Biol 10:1048–53
Paterson RG, Russell CJ, and Lamb RA (2000) Fusion protein of the paramyxovirus SV5: destabilizing and stabilizing mutants of fusion activation. Virology 270:17–30
Pelkmans L and Helenius A (2003) Insider information: what viruses tell us about endocytosis. Curr Opin Cell Biol 15:414–22
Percherancier Y, Lagane B, Planchenault T, Staropoli I, Altmeyer R, Virelizier JL, Arenzana-Seisdedos F, Hoessli DC, and Bachelier F (2003) HIV-1 entry into T-cells is not dependent on CD4 and CCR5 localization to sphingolipid-enriched, detergent-resistant, raft membrane domains. J Biol Chem 278:3153–3161
Pereira FB, Goni FM, and Nieva JL (1995) Liposome destabilization induced by the HIV-1 fusion peptide effect of a single amino acid substitution. FEBS Lett 362:243–6
Pietschmann T, Zentgraf H, Rethwilm A, and Lindemann D (2000) An evolutionarily conserved positively charged amino acid in the putative membrane-spanning domain of the foamy virus envelope protein controls fusion activity. J Virol 74:4474–82
Piller SC, Dubay JW, Derdeyn CA, and Hunter E (2000) Mutational analysis of conserved domains within the cytoplasmic tail of gp41 from human immunodeficiency virus type 1: effects on glycoprotein incorporation and infectivity. J Virol 74:11717–23
Pinter A, Kopelman R, Li Z, Kayman SC, and Sanders DA (1997) Localization of the labile disulfide bond between SU and TM of the murine leukemia virus envelope protein complex to a highly conserved CWLC motif in SU that resembles the active-site sequence of thiol-disulfide exchange enzymes. J Virol 71:8073–7
Popik W, Alce TM, and Au WC (2002) Human immunodeficiency virus type 1 uses lipid raft-coloalized CD4 and chemokine receptors for productive entry into CD4(+) T cells. J Virol 76:4709–22
Puri A, Winick J, Lowy RJ, Covell D, Eidelman O, Walter A, and Blumenthal R (1988) Activation of vesicular stomatitis virus fusion with cells by pretreatment at low pH. J Biol Chem 263:4749–53
Qiao H, Armstrong RT, Melikyan GB, Cohen FS, and White JM (1999) A specific point mutant at position 1 of the influenza hemagglutinin fusion peptide displays a hemifusion phenotype. Mol Biol Cell 10:2759–69
Qiao H, Pelletier SL, Hoffman L, Hacker J, Armstrong RT, and White JM (1998) Specific single or double proline substitutions in the “spring-loaded” coiled-coil region of the influenza hemagglutinin impair or abolish membrane fusion activity. J Cell Biol 141:1335–47

Ragheb JA and Anderson WF (1994) pH-independent murine leukemia virus ecotropic envelope-mediated cell fusion: implications for the role of the R peptide and p12E TM in viral entry. J Virol 68:3220–31

Reeves JD, Gallo SA, Ahmad N, Miamidian JL, Harvey PE, Sharron M, Pohlman S, Sfakianos JN, Derdeyn CA, Blumenthal R, Hunter E, and Doms RW (2002) Sensitivity of HIV-1 to entry inhibitors correlates with envelope/coreceptor affinity, receptor density, and fusion kinetics. Proc Natl Acad Sci USA 99:16249–54

Rein A, Mirro J, Haynes JG, Ernst SM, and Nagashima K (1994) Function of the cytoplasmic domain of a retroviral transmembrane protein: p15E-p2E cleavage activates the membrane fusion capability of the murine leukemia virus Env protein. J Virol 68:1773–81

Rey FA, Heinz FX, Mandl C, Kunz C, and Harrison SC (1995) The envelope glycoprotein from tick-borne encephalitis virus at 2 Å resolution. Nature 375:291–8

Roussko I, Mixon MB, Chen BK, and Kim PS (2000) Palmitoylation of the HIV-1 envelope glycoprotein is critical for viral infectivity. Proc Natl Acad Sci USA 97:13523–5

Russell CJ, Jardetzky TS, and Lamb RA (2001) Membrane fusion machines of paramyxoviruses: capture of intermediates of fusion. EMBO J 20:4024–34

Saez-Cirion A, Gomara MJ, Agirre A, and Nieva JL (2003) Pre-transmembrane sequence of Ebola glycoprotein. Interfacial hydrophobicity distribution and interaction with membranes. FEBS Lett 533:47–53

Saez-Cirion A, Nir S, Lorizate M, Agirre A, Cruz A, Perez-Gil J, and Nieva JL (2002) Sphingomyelin and cholesterol promote HIV-1 gp41 pretransmembrane sequence surface aggregation and membrane restructuring. J Biol Chem 277:21776–85

Sakai T, Ohuchi R, and Ohuchi M (2002) Fatty acids on the A/USSR/77 influenza virus hemagglutinin facilitate the transition from hemifusion to fusion pore formation. J Virol 76:4603–11

Salzwedel K, West JT, and Hunter E (1999) A conserved tryptophan-rich motif in the membrane-proximal region of the human immunodeficiency virus type 1 gp41 ectodomain is important for Env-mediated fusion and virus infectivity. J Virol 73:2469-80

Schibli DJ, Montelaro RC, and Vogel HJ (2001) The membrane-proximal tryptophan-rich region of the HIV glycoprotein, gp41, forms a well-defined helix in dodecylphosphocholine micelles. Biochemistry 40:9570–8

Schmid E, Zurbriggen A, Gassen U, Rima B, ter Meulen V, and Schneider-Schaufes J (2000) Antibodies to CD9, a tetraspan transmembrane protein, inhibit canine distemper virus-induced cell-cell fusion but not virus-cell fusion. J Virol 74:7554–61

Schoch C and Blumenthal R (1993) Role of the fusion peptide sequence in initial stages of influenza hemagglutinin-induced cell fusion. J Biol Chem 268:9267–74

Schultz A and Rein A (1985) Maturation of murine leukemia virus Env proteins in the absence of other viral proteins. Virology 145:335–9
Seth S, Vincent A, and Compans RW (2003) Mutations in the cytoplasmic domain of a paramyxovirus fusion glycoprotein rescue syncytium formation and eliminate the hemagglutinin-neuraminidase protein requirement for membrane fusion. J Virol 77:167-78
Shin JS and Abraham SN (2001) Caveolae as portals of entry for microbes. Microbes Infect 3:755–61
Sjoberg M and Garoff H (2003) Interactions between the transmembrane segments of the alphavirus E1 and E2 proteins play a role in virus budding and fusion. J Virol 77:3441–3450
Skehel JJ, Cross K, Steinhauer D, and Wiley DC (2001) Influenza fusion peptides. Biochem Soc Trans 29:623–6
Skehel JJ and Wiley DC (2000) Receptor binding and membrane fusion in virus entry: the influenza hemagglutinin. Annu Rev Biochem 69:531–69
Staschke KA, Hatch SD, Tang JC, Hornback WJ, Munroe JE, Colacino JM, and Muesing MA (1998) Inhibition of influenza virus hemagglutinin-mediated membrane fusion by a compound related to podocarpic acid. Virology 248:264–274
Stegmann T, Bartoldus I, and Zumbrunn J (1995) Influenza hemagglutinin-mediated membrane fusion: influence of receptor binding on the lag phase preceding fusion. Biochemistry 34:1825–32
Stegmann T, Booy FP, and Wilschut J (1987) Effects of low pH on influenza virus. Activation and inactivation of the membrane fusion capacity of the hemagglutinin. J Biol Chem 262:17744–9
Stein BS, Gowda SD, Lifson JD, Penhallow RC, Bensch KG, and Engleman EG (1987) pH-independent HIV entry into CD4-positive T cells via virus envelope fusion to the plasma membrane. Cell 49:659–68
Stiasny K, Allison SL, Mandl CW, and Heinz FX (2001) Role of metastability and acidic pH in membrane fusion by tick-borne encephalitis virus. J Virol 75:7392–7398
Stiasny K, Allison SL, Marchler-Bauer A, Kunz C, and Heinz FX (1996) Structural requirements for low-pH-induced rearrangements in the envelope glycoprotein of tick-borne encephalitis virus. J Virol 70:8142–8147
Stiasny K, Allison SL, Schalich J, and Heinz FX (2002) Membrane interactions of the tick-borne encephalitis virus fusion protein E at low pH. J Virol 76:3784–90
Stone-Hulslander J and Morrison TG (1997) Detection of an interaction between the HN and F proteins in Newcastle disease virus-infected cells. J Virol 71:6287–95
Suarez T, Gallaher WR, Agirre A, Goni FM, and Nieva JL (2000) Membrane interface-interacting sequences within the ectodomain of the human immunodeficiency virus type 1 envelope glycoprotein: putative role during viral fusion. J Virol 74:8038–47
Suomalainen M (2002) Lipid rafts and assembly of enveloped viruses. Traffic 3:705–9
Taguchi F and Matsuyama S (2002) Soluble receptor potentiates receptor-independent infection by murine coronavirus. J Virol 76:950–8
Takimoto T, Taylor GL, Connaris HC, Crennell SJ, and Portner A (2002) Role of the hemagglutinin-neuraminidase protein in the mechanism of paramyxovirus-cell membrane fusion. J Virol 76:13028–33
Tamm LK and Han X (2000) Viral fusion peptides: a tool set to disrupt and connect biological membranes. Biosci Rep 20:501–18
Tamm LK, Han X, Li Y, and Lai AL (2002) Structure and function of membrane fusion peptides. Biopolymers 66:249–60
Taylor GM and Sanders DA (1999) The role of the membrane-spanning domain sequence in glycoprotein-mediated membrane fusion. Mol Biol Cell 10:2803–15
Tong S and Compans RW (2000) Oligomerization, secretion, and biological function of an anchor-free parainfluenza virus type 2 (PI2) fusion protein. Virology 270:368–76
Tong S, Li M, Vincent A, Compans RW, Fritsch E, Beier R, Klenk C, Ohuchi M, and Klenk HD (2002) Regulation of fusion activity by the cytoplasmic domain of a paramyxovirus F protein. Virology 301:322–333
Vashishtha M, Phalen T, Marquardt MT, Ryu JS, Ng AC, and Kielen M (1998) A single point mutation controls the cholesterol dependence of Semliki Forest virus entry and exit. J Cell Biol 140:91–9
Viard M, Parolini I, Sargiacomo M, Fecchi K, Ramoni C, Ablan S, Ruscetti FW, Wang JM, and Blumenthal R (2002) Role of cholesterol in human immunodeficiency virus type 1 envelope protein-mediated fusion with host cells. J Virol 76:11584–95
Waarts BL, Bittman R, and Wilschut J (2002) Sphingolipid and cholesterol dependence of alphavirus membrane fusion. Lack of correlation with lipid raft formation in target liposomes. J Biol Chem 277:38141–38147
Wahlberg JM, Bron R, Wilschut J, and Garoff H (1992) Membrane fusion of Semliki Forest virus involves homotrimers of the fusion protein. J Virol 66:7309–18
Wahlberg JM and Garoff H (1992) Membrane fusion process of Semliki Forest virus. I: Low pH-induced rearrangement in spike protein quaternary structure precedes virus penetration into cells. J Cell Biol 116:339–48
Weber T, Paesold G, Galli C, Mischler R, Semenza G, and Brunner J (1994) Evidence for H+-induced insertion of influenza hemagglutinin HA2 N-terminal segment into viral membrane. J Biol Chem 269:18353–8
Weissenhorn W, Carfi A, Lee KH, Skehel JJ, and Wiley DC (1998) Crystal structure of the Ebola virus membrane fusion subunit, GP2, from the envelope glycoprotein ectodomain. Mol Cell 2:605–16
Weissenhorn W, Dessen A, Calder LJ, Harrison SC, Skehel JJ, and Wiley DC (1999) Structural basis for membrane fusion by enveloped viruses. Mol Membr Biol 16:3–9
Weissenhorn W, Dessen A, Harrison SC, Skehel JJ, and Wiley DC (1997) Atomic structure of the ectodomain from HIV-1 gp41. Nature 387:426–30
West JT, Johnston PB, Dubay SR, and Hunter E (2001) Mutations within the putative membrane-spanning domain of the simian immunodeficiency virus transmembrane glycoprotein define the minimal requirements for fusion, incorporation, and infectivity. J Virol 75:9601–12
Wharton SA, Skehel JJ, and Wiley DC (2000) Temperature dependence of fusion by Sendai virus. Virology 271:71–78
White J and Helenius A (1980) pH-dependent fusion between the Semliki Forest virus membrane and liposomes. Proc Natl Acad Sci USA 77:3273–7
White J, Kartenbeck J, and Helenius A (1980) Fusion of Semliki forest virus with the plasma membrane can be induced by low pH. J Cell Biol 87:264–72
White J, Kartenbeck J, and Helenius A (1982) Membrane fusion activity of influenza virus. EMBO J 1:217–22
White JM (1990) Viral and cellular membrane fusion proteins. Annu Rev Physiol 52:675–97
Wilson IA, Skehel JJ, and Wiley DC (1981) Structure of the haemagglutinin membrane glycoprotein of influenza virus at 3 Å resolution. Nature 289:366–373
Xiang SH, Kwong PD, Gupta R, Rizzuto CD, Casper DJ, Wyatt R, Wang L, Hendrickson WA, Doyle ML, and Sodroski J (2002) Mutagenic stabilization and/or disruption of a CD4-bound state reveals distinct conformations of the human immunodeficiency virus type 1 gp120 envelope glycoprotein. J Virol 76:9888–9899
Yang C and Compans RW (1996) Analysis of the cell fusion activities of chimeric simian immunodeficiency virus-murine leukemia virus envelope proteins: inhibitory effects of the R peptide. J Virol 70:248–54
Yang L and Huang HW (2002) Observation of a membrane fusion intermediate structure. Science 297:1877–9
Yao Q, Hu X, and Compans RW (1997) Association of the parainfluenza virus fusion and hemagglutinin-neuraminidase glycoproteins on cell surfaces. J Virol 71:650–6
Young JK, Li D, Abramowitz MC, and Morrison TG (1999) Interaction of peptides with sequences from the Newcastle disease virus fusion protein heptad repeat regions. J Virol 73:5945–56
Zarkik S, Defrise-Quertain F, Portetelle D, Burny A, and Ruysschaert JM (1997) Fusion of bovine leukemia virus with target cells monitored by R18 fluorescence and PCR assays. J Virol 71:738–40
Zhou J, Dutch RE, and Lamb RA (1997) Proper spacing between heptad repeat B and the transmembrane domain boundary of the paramyxovirus SV5 F protein is critical for biological activity. Virology 239:327–39