Enveloped viruses initiate infection by fusion of the viral membrane with a membrane of the presumptive host cell. Conformational changes in surface-expressed, membrane-anchored “fusion proteins,” coupled with attachment to the target membrane, overcome the kinetic barrier to bilayer merger (1, 2). A general model for these fusion-inducing conformational changes, derived from studies of many viral fusion proteins, invokes a canonical sequence of events: a priming step, often a proteolytic cleavage and usually irreversible; a triggering step, such as exposure to low pH in endosomes or, for some viruses, receptor binding; formation of an extended intermediate, from which hydrophobic fusion loops or fusion peptides insert into the target membrane; and collapse of that intermediate to a final, stable conformation that brings together the fusion loops or peptides and the transmembrane anchor, and hence pulls together the two membranes (3). Structures of the initial (prefusion) conformation, both unprimed and primed, and the final (postfusion) conformation have shown the beginning and end of the fusion process for many enveloped viruses (4); studies of single virus–particle fusion kinetics have probed the intervening stages in some detail for influenza and West Nile viruses (5–7).

The fusion glycoprotein (G protein) of vesicular stomatitis virus (VSV) and related rhabdoviruses (e.g., rabies virus) is the sole surface-expressed protein on the bullet-shaped virions. It mediates both attachment and low-pH-induced fusion (8). Its fusogenic conformational changes deviate from the canonical sequence outlined in the preceding paragraph by the absence of an irreversible priming step and hence the absence of a metastable prefusion state. The transition from prefusion conformation to extended intermediate is reversible (9, 10). Nonetheless, structures of G in its pre- and postfusion trimeric conformations suggest that most of the fusion reaction follows a familiar pattern, as illustrated in Fig. 1 (3, 11–13). We show the extended intermediate as a monomer, because the two structures appear to require a dissociative transition from pre- to postfusion trimer (Fig. 1, open and extended conformations). Note that in this inferred picture of the transition from prefusion to postfusion conformations, the exposed lateral surfaces of the apical domain of the molecule (those facing left and right in the first panel of Fig. 1) become buried along the threefold contact when the extended intermediate trimerizes and that the extended C-terminal segment “zips” up along the outside of this trimer during the fold-back step.

We report here single-virion fusion experiments, carried out on VSV. Fusion of two lipid bilayers generally proceeds through a hemifusion state, in which the apposed leaflets have merged but not the distal leaflets (6); we can detect hemifusion by observing transfer of a fluorescent molecule from one membrane to the other. In particular, we have inserted a lipophilic dye, R18,
into the viral membrane at self-quenching concentrations and recorded the time elapsed between a fusion-inducing pH drop and dequenching of the fluorophore followed by dissipation of its fluorescent signal by diffusion in the target membrane. We find that the kinetic data are indeed consistent with the general picture shown in Fig. 1. By varying both initial and final pH, we can separate the kinetic steps and show that the following three-stage model fits the observations. The first step is a reversible, pH-dependent G-protein conformational change, corresponding to the transition from “prefusion” to “extended” in Fig. 1; the second, a reversible G trimerization and clustering of fusion loops (“extended” to “trimerized”); and the third, folding back to bring together the two membranes. The membranes resist this collapse, and a critical number of adjacent, extended trimers spanning the contact zone between virus and target membrane are necessary to progress forward to hemifusion. Computational simulations match the observations if the critical cluster is chosen as 3, 4, or 5 trimers, within a contact zone of 30 to 50 G trimers. The time required to accumulate this critical cluster determines the overall rate of the fusion reaction. This mechanism is essentially the same as the ones previously described for influenza and West Nile viruses (5–7), despite differences in the structures of their fusion proteins, which represent each of the three “classes” so far described (3).

**Results**

**pH Dependence of VSV Hemifusion.** We used total internal reflection fluorescence (TIRF) microscopy, in a configuration similar to the one described previously (Fig. 2 A and B) (6), to follow single-particle VSV fusion with a supported lipid bilayer under nine different regimes of initial and final pH (Fig. 2 C–F). We labeled the VSV membrane with quenching concentrations of a lipophilic dye, R18. The virus attached to the bilayer by the initial pH; a rapid pH drop initiated the fusogenic conformational change in G. At the time of hemifusion, R18 could diffuse from the viral membrane into the target bilayer, leading to dequenching of the R18 signal and then rapid dissipation (Fig. 2B and Movie S1). For each experiment, the initial pH was higher than that of the fusion threshold (pH 6.4) (14). The mean time from pH drop (detected by the pH-dependent loss of fluorescence from the fluorescein incorporated into the membrane) to hemifusion (detected by R18 dequenching) depended on both initial and final pH (Fig. 2C); it decreased as either limiting pH decreased. At the lowest final pH (5.5), the mean time to hemifusion approached 25 to 30 s, regardless of the initial pH.

At high initial or final pH, the hemifusion frequency distributions showed a rise and decay, the signature of more than one rate-limiting step (Fig. 2 D and E, Top). As the final pH decreased, the distribution shifted to an exponential decay (Fig. 2 D–F, Bottom), the characteristic distribution for a single rate-limiting step. A similar trend applied to the initial pH (Fig. 2 D–F, Top) but, even at pH 6.6, the shape of the distribution was not a simple exponential.

**pH Dependence of VSV–Membrane Association.** During the hemifusion experiments, we observed that the VSV particle attached to the membrane in two modes. In a “rolling” mode, the virions moved along the bilayer in the direction of flow, while clearly maintaining contact with the bilayer, as they remained within the TIRF evanescent field (Fig. 3A, particle 1). In an “arrested” mode, virions were immobile, even if subject to drag in the flow cell of the microscope (Fig. 3A, particles 2 and 3; see also Movie S2). Both rolling and arrested virions underwent hemifusion following the drop in pH. Rolling virions arrested rapidly during the short (2- to 4-s) period that marked the transition from initial to final pH, and all particles had arrested by the time, t0, at which the pH in the flow cell had dropped to its final level. When the initial pH was 6.6, all virions were arrested, even before the pH drop. Because pH 6.6 is also the point at which a single, rate-limiting step determines the hemifusion time distribution and other steps become much faster, this result suggests that the molecular transitions responsible for virion rolling and arrest are related to the pH-dependent steps in VSV hemifusion.

Fig. 3B shows a more detailed analysis of the pH dependence of rolling. Virions bound to the target bilayer were equilibrated for 5 to 10 min at one pH and then imaged under flow at that same pH. The percentage of rolling virions decreased roughly linearly with pH (Fig. 3B). At pH 8.0, over 60% of the bound virions were moving; at pH 7.4, about 30%; at pH 6.6, virtually all bound virions were stationary, consistent with our observations in the pH-drop experiments. The mean speed of the rolling population of virions also decreased steadily with pH (Fig. 3C).

The transition between rolling and arrest is reversible. When we raised the pH from 6.6, at which all virions were stationary, to 7.4, many of the arrested virions began to roll. When the pH was
lowered again to 6.6, the rolling stopped (Fig. 3D). We interpret this observation by assuming a reversible, pH-dependent transition from the prefusion conformation, in which the fusion loops of G project back toward the viral membrane around the periphery of the trimer (Fig. 1, prefusion conformation), to an extensible monomer, in which the fusion loops can contact the target membrane (Fig. 1, extended conformation). Because we did not use a surrogate receptor in our experiments [such as the equivalent of a ganglioside in work on influenza virus fusion (6, 7) or a lectin domain in studies of West Nile virus fusion (5)], attachment to the supported bilayer was probably through irreversible exposure of the fusion loops, even at pH 8.0. As the pH dropped, the equilibrium shifted toward loop exposure, and more extensive interactions anchored the particle firmly enough to resist solvent drag. Reversibility of the rolling phenomenon indicates that membrane interaction of extended G at pH ≥6.6 is itself reversible.

**pH Transitions and Conformation of VSV G.** The pH dependence of virion binding derives from conformational changes in the surface-expressed G protein. We examined changes in the G-protein layer by negative-stain electron microscopy of VSV particles incubated at pH 7.6, 7.0, and 6.6 (Fig. S1). As previously reported (15), the G layer at pH 7.6 was shallow (average depth 6.0 ± 0.4 nm) and appeared indistinct (Fig. S1A), whereas at pH 6.6, most of the G layer was deeper (average depth 10.5 ± 0.6 nm) and appeared more ordered (Fig. S1C). At pH 7.0, patches of the longer form of G appeared interspersed with patches of the shorter form of G (Fig. S1B). These observations suggest that as the pH decreases from 7.6 to 6.6, the G layer of the particle gradually converts from the short form to the long form.

We used a liposome-binding experiment to estimate the pK<sub>a</sub> of the transition of G into a membrane-interacting conformation. We generated the G ectodomain (G<sub>ex</sub>) by thermolysin cleavage of intact virus particles and purified it by anion-exchange chromatography. We also made G<sub>ex</sub>-W72A, both wild-type (G<sub>ex</sub>-WT) and mutant (G<sub>ex</sub>-W72A), with liposomes at several pH values, separated the liposome-bound from free protein by sucrose-density centrifugation, and detected G<sub>ex</sub> in each fraction of the gradient by immunoblotting with a conformation-specific monoclonal antibody, IE2 (17).

At pH 8.0 and 7.4, most of the G<sub>ex</sub>-WT remained at the bottom of the gradient; at pH 6.6 and 6.0, most of it shifted to the top of the gradient, showing association (“cofлотation”) with the liposomes in that fraction (Fig. 4B). G<sub>ex</sub>-W72A did not associate with liposomes at any pH (Fig. 4C). The transition between pH 7.4 and 6.6 corresponds closely to the transition between rolling and arrest. Moreover, both transitions are reversible: Back-neutralization to pH 8.0 of G<sub>ex</sub>-WT incubated with liposomes at

![Fig. 2. Hemifusion kinetics of VSV. (A) Representative TIRF image of labeled virus particles bound to a lipid bilayer. Virus particles, whose membranes are labeled with fluorescent lipophilic dye R18, enter the flow-cell channel and bind nonspecifically to a target lipid bilayer, supported on a glass coverslip. Fluoresceinated lipid in the bilayer acts as an internal pH sensor, as the fluorescence yield of fluorescein decreases with pH. Introduction of low-pH buffer triggers hemifusion. (B) Snapshots of a single hemifusing virus particle (Top) and corresponding intensity traces (Bottom). The fluorescein intensity trace is in green; the R18 intensity trace, in red; and the pH sensor trace, in blue. Hemifusion is detected by a sharp rise in intensity due to dequenching of the lipophilic R18 dye upon lipid mixing. (C) Mean hemifusion times measured at varied initial and final pH values. VSV particles bound to the target bilayer were incubated at one of three initial pH values (legend). The pH was then lowered to one of three final pH values (x axis) to initiate hemifusion. The mean hemifusion times were calculated from the hemifusion time distributions shown in D–F. Error bars show standard deviation (SD). (D–F) Distributions of hemifusion times measured at varied initial and final pH values. Initial pH at the top of each figure; final pH, within each plot. Each vertical bind represents the fraction of the VSV population that has hemifused within the time interval indicated on the x axis. The number of bins in each histogram does not exceed the square root of the number of virions in each distribution. The heights of the bins have been normalized such that the area under the curve equals one.

![Fig. S1.](https://example.com/FigS1.png)
pH 6.6 or 6.0 eliminated the coflotation (Fig. 4D), just as back-neutralization restored rolling. The failure of G<sub>WT</sub>-W72A to comigrate with liposomes on the gradient at any pH confirms that the properties of G<sub>WT</sub> probed in these experiments are indeed due to fusion-loop interactions with the lipid bilayer.

Discussion

Structural Interpretation of Hemifusion Kinetics. The soluble ectodomain, G<sub>WT</sub>, has the following conformational properties. At pH > 7, it is in equilibrium between the “umbrella-like” prefusion trimer seen in crystals grown at elevated pH and an extended monomer. The interactions among subunits in the prefusion trimer are weak enough that the soluble ectodomain is monomeric at concentrations (~1 ng/mL) ordinarily used for biochemical characterization (18). At pH < 7 and in the presence of a lipid bilayer (e.g., the liposomes in the experiments shown in Fig. 4), membrane binding through the fusion loops will favor trimer clustering and folding back into the inverted, postfusion conformation seen in crystals at acidic pH. Stable association with liposomes (and hence detectable coflotation) probably requires the joint participation of all three subunits. The conformational change and liposome binding are nonetheless reversible upon reneutralization (Fig. 4D), and the individual extended monomers can dissociate from the liposome; at suitable concentrations, they will also reform the soluble, prefusion trimer. Soluble forms of flavivirus E proteins show a similar, liposome-catalyzed trimerization, but in that case an irreversible one (19).

On the surface of a virion at neutral pH and above, our results together with published data indicate that full-length G is in equilibrium between the prefusion trimer conformation and flexibly extended monomers (10, 20–22). In the absence of a target membrane, irreversible transition of virion G to its conformation at the end of a complete fusion reaction would require that three subunits come together, fold back, and insert their fusion loops into the viral membrane (Fig. 1). There is a barrier to this transition even at pH 6.6, however, because exposure to that pH does not inactivate the virus (and because, in the presence of a target membrane, progression to hemifusion and fusion is immeasurably slow). Proton binding at pH ∼ 6.4 and below

---

**Fig. 3.** VSV particles bind the target membrane in two modes. (A) Successive frames of a movie showing the two modes of binding at pH 7.4. Three particles are shown in white. Particles 2 and 3 are arrested and particle 1 rolls. Each frame is a 400-ms exposure taken every 2 s. The flow direction is down. (B) Percentage of VSV particles that are rolling at pH values between 8.0 and 6.6. Each point represents the average of three to four independent trials. Error bars show SD. (C) Distributions of mean velocities of rolling VSV particles at pH values between 8.0 and 6.6. Each bin represents the number of particles rolling with a mean velocity within the values indicated on the x axis. (D) Successive frames of a movie showing the reversibility of the transition between rolling and arrest. The particle is shown in white. The pH value at each time is indicated above the frames. Frames are 300-ms exposures taken every 3 s. The flow direction is down. The images shown in A and D have been sharpened with a Mexican hat filter and their contrast increased to aid in particle tracking.

---

**Fig. 4.** Liposome interaction of a G fusion-loop mutant. (A) Close-up view of conserved hydrophobic residues at the tips of each fusion loop in a G trimer. Each G monomer is colored differently. The conserved hydrophobic residues—W72, Y73, Y116, and A117—are shown as sticks [PDB ID code 5I2M (11)]. (B and C) Cleaved ectodomain (G<sub>T</sub>) from G<sub>WT</sub> (B) or the fusion-loop mutant, G-W72A (C), was incubated with liposomes at the listed pH values. The mixtures were then separated over a discontinuous sucrose gradient, and each fraction was immunoblotted with the monoclonal antibody IE2 to detect the presence of G<sub>E</sub>. Fraction numbers are along the bottom of each figure; the top (T) and bottom (B) fractions are also labeled. (D) In the reversibility experiment, G<sub>T</sub>-WT was initially incubated with liposomes at the indicated starting pH before the pH was shifted to 8.0. After further incubation at pH 8.0, the mixtures were separated over a discontinuous sucrose gradient, also at pH 8.0.
Buffers used for virus purification were HNE-10 pH 8.0 (50 mM Hepes, pH 8.0, 130 mM NaCl, 0.1 mM EDTA), HNE-50 pH 7.4 (50 mM Hepes, pH 7.4, 140 mM NaCl, 0.1 mM EDTA), and MES-50 pH 6.6 (50 mM MES, pH 6.6, 130 mM NaCl, 0.1 mM EDTA). The final pH buffers used were HNE-50 pH 8.0 (50 mM Hepes, pH 8.0, 130 mM NaCl, 0.1 mM EDTA), HNE-50 pH 7.4 (50 mM Hepes, pH 7.4, 140 mM NaCl, 0.1 mM EDTA), and MES-50 pH 6.6 (50 mM MES, pH 6.6, 130 mM NaCl, 0.1 mM EDTA). The increase in buffer concentration sharpened the transition from the initial pH to the final pH during the experiment. The sodium chloride concentration in these buffers was adjusted such that the total ionic strength of the buffer was ∼150 mM. Ionic strength was calculated using the formula given in ref. 26. The initial pH buffers used were HNE-50 pH 8.0 (50 mM Hepes, pH 8.0, 130 mM NaCl, 0.1 mM EDTA), HNE-50 pH 7.4 (50 mM Hepes, pH 7.4, 140 mM NaCl, 0.1 mM EDTA), and MES-50 pH 6.6 (50 mM MES, pH 6.6, 130 mM NaCl, 0.1 mM EDTA). The final pH buffers used were MES-100 pH 6.2 (100 mM MES, pH 6.2, 122 mM NaCl, 0.1 mM EDTA), MES-100 pH 6.0 (100 mM MES, pH 6.0, 128 mM NaCl, 0.1 mM EDTA), and MES-100 pH 5.5 (100 mM MES, pH 5.5, 140 mM NaCl, 0.1 mM EDTA).

Materials and Methods

Buffer Solutions. Buffers used for virus purification were HNE-10 pH 8.0 (10 mM Hepes, pH 8.0, 140 mM NaCl, 0.1 mM EDTA) and HNE-10 pH 7.4 (10 mM Hepes, pH 7.4, 140 mM NaCl, 0.1 mM EDTA).

In the hemifusion experiments, buffers at the initial pH values contained either 50 mM Hepes or 50 mM MES and buffers at the final pH values contained 100 mM MES. The increase in buffer concentration sharpened the transition from the initial pH to the final pH during the experiment. The sodium chloride concentration in these buffers was adjusted such that the total ionic strength of the buffer was ∼150 mM. Ionic strength was calculated using the formula given in ref. 26. The initial pH buffers used were HNE-50 pH 8.0 (50 mM Hepes, pH 8.0, 130 mM NaCl, 0.1 mM EDTA), HNE-50 pH 7.4 (50 mM Hepes, pH 7.4, 140 mM NaCl, 0.1 mM EDTA), and MES-50 pH 6.6 (50 mM MES, pH 6.6, 130 mM NaCl, 0.1 mM EDTA). The final pH buffers used were MES-100 pH 6.2 (100 mM MES, pH 6.2, 122 mM NaCl, 0.1 mM EDTA), MES-100 pH 6.0 (100 mM MES, pH 6.0, 128 mM NaCl, 0.1 mM EDTA), and MES-100 pH 5.5 (100 mM MES, pH 5.5, 140 mM NaCl, 0.1 mM EDTA).

In the rolling experiments, the buffers used were HNE-50 pH 8.0 (50 mM Hepes, pH 8.0, 130 mM NaCl, 0.1 mM EDTA), HNE-50 pH 7.6 (50 mM Hepes, pH 7.6, 137 mM NaCl, 0.1 mM EDTA), HNE-50 pH 7.4 (50 mM Hepes, pH 7.4, 140 mM NaCl, 0.1 mM EDTA), HNE-50 pH 7.0 (50 mM Hepes, pH 7.0, 147 mM NaCl, 0.1 mM EDTA), HNE-50 pH 6.6 (50 mM Hepes, pH 6.6, 147 mM NaCl, 0.1 mM EDTA), MES-100 pH 6.8 (100 mM MES, pH 6.8, 108 mM NaCl, 0.1 mM EDTA), and MES-100 pH 6.6 (100 mM MES, pH 6.6, 112 mM NaCl, 0.1 mM EDTA).

All buffers used in imaging experiments were supplemented with 1 mM CaCl₂, 1 mM MgCl₂, and an oxygen-scavenging system composed of protoocatechuate 3,4-dioxygenase from Pseudomonas (PCD; Sigma-Aldrich), 3,4-dihydroxybenzoic acid (protocatechuic acid; PCA; Sigma-Aldrich), and (±)-6-hydroxy-2,5,7,8-tetramethylchroman-2-carboxylic acid (Trolox;Sigma-Aldrich) at final concentrations of 100 mM, 2.5 mM, and 1 mM, respectively. The components of the oxygen-scavenging system were prepared as described (27).

Cells. BSR-T7 cells (28) and Vero cells (ATCC) were grown at 37 °C in Dulbecco’s modified Eagle medium (DMEM; Gibco) supplemented with 10% (vol/vol) FBS (Gibco).

Virus Growth and Purification. Recombinant vesicular stomatitis virus (rVSV) was propagated in BSR-T7 cells and purified as follows (29). BSR-T7 cell monolayers were inoculated with rVSV (multiplicity of infection of 0.1) in DMEM supplemented with 2% (vol/vol) FBS and the antibiotics penicillin, streptomycin, and kanamycin for 18 to 20 h at 34 °C. After a low-speed centrifugation step to clear cell debris (2,000 × g for 5 min), virus particles were pelleted from the medium of infected cells by ultracentrifugation at 17,000 rpm for 1.5 h at 4 °C in an SW28 rotor (Beckman Coulter) and resuspended in...
either HNE-10 pH 8.0 or HNE-10 pH 7.4 (Buffer Solutions) overnight at 4 °C. The concentrated virus suspension was further separated on a linear 15 to 45% (wt/vol) sucrose gradient formed in either HNE-10 pH 8.0 or HNE-10 pH 7.4 at 25,000 rpm for 3.5 h at 4 °C in an SW41 rotor (Beckman Coulter). The bottom band of virus particles was harvested by side puncture and concentrated by centrifugation through a 10% (wt/vol) sucrose cushion in either HNE-10 pH 8.0 or HNE-10 pH 7.4 at 33,000 rpm for 1 h at 4 °C in an SW50.1 rotor (Beckman Coulter). The virus pellet was resuspended in either HNE-10 pH 8.0 or HNE-10 pH 7.4 overnight at 4 °C. Purity and protein content of the virus particles were determined by SDS-PAGE and Coomassie staining. Viral titers were measured on monolayers of Vero cells as previously described (30).

Fig. 5. Kinetic model for VSV fusion. (A1) Monomer extension (reversible). On the virion surface, we assume that G forms loosely associated prefusion trimers, in which each monomer is in the prefusion form G0. Each G0 monomer can protonate independently and extend reversibly to form extended monomer, G*, exposing the fusion loops. The relative concentrations of G0 and G* at equilibrium are determined by the pH and an apparent pK1a. The forward rate constant for extension is k1, and the reverse rate constant is k−1, or k1/(10^−pK1a). (A2) Trimerization (reversible). Three adjacent extended G* monomers reversibly associate to form an extended trimer, G3. The forward and reverse rate constants of this trimerization are k2 and k2−, respectively. To simplify the model, we assume that G* only trimerizes with its original prefusion trimer neighbors. (A3) Hemifusion (irreversible). A cluster of extended G3 trimers folds back to mediate hemifusion. The number of trimers required to form the fold-back cluster is n. The irreversible fold-back step proceeds with rate constant k3. In this diagram, a cluster of size n = 4 is shown. G proteins elsewhere in the particle can independently adopt any of the previously described conformations of G. (B) Fits of the model to experimental data (see also Materials and Methods). Shown here is the best fit (black line) obtained for patch size P = 55 trimers and fold-back cluster size n = 4, where pK1a = 7.1, k1 = 9.1 × 10^5 mol⁻¹ L s⁻¹, k2 = 2.0 s⁻¹, k2− = 9.9 s⁻¹, and k3 = 5.8 s⁻¹. Experimental data (red bars) are from Fig. 2 D–F.
The total protein concentration of purified VSV was determined by Bradford assay (Bio-Rad) using a BSA standard. To label the virus with lipophilic dye, 50 to 100 μL purified VSV (1 mg/mL in HNE-50 pH 8.0 or HNE-50 pH 7.4) was mixed with 0.5 to 1 μL octadecyl rhodamine B chloride (R18; 2 mM in ethanol; Invitrogen) for a final R18 concentration of 20 μM and incubated at room temperature (RT) for 1.5 to 2 h. The labeled virus was separated from unincorporated dye over a gel-filtration column (PD-10 desalting column or PD MidiTrap G-25 desalting column; GE Healthcare). There was less dilution of the labeled virus fraction when using the PD MidiTrap G-25 desalting column.

**Flow-Cell Construction.** Glass microscope coverslips (25 × 25 mm; no. 1.0; VWR) were imaged by serial rounds of sonication in 7X-O-Matic detergent (VWR), 1 M potassium hydroxide, HPLC-grade acetone, and HPLC-grade ethanol for 10 to 20 min each round. The glass coverslips were thoroughly rinsed in Milli-Q water (Millipore) after each round of sonication. After the final sonication in ethanol, the coverslips were dried by baking at 110 °C. To render the surface of the glass hydrophilic, the coverslips were cleaned with oxygen plasma (0.5 torr) for 3 min (plasma etcher; March Plasma). The flow cell was constructed by placing a poly(dimethylsiloxane) (PDMS) device containing microfluidic channels (70 μm high × 0.5 mm wide × ~5 mm long) onto a coverslip and sealing by compression. Inlet and outlet tubing was connected through holes bored in the PDMS device. To minimize the dead volume, inlet tubing with an inner diameter of 200 μm (Teflon FEP tubing; IDEX Health & Science) and a length of 6 cm was used. The outlet tubing (PE60 tubing, inner diameter 0.38 mm; BD Biosciences) was connected to a syringe pump (Harvard Apparatus), and flow was established by negative pressure across the channel.

**Lipid Bilayer Preparation.** Liposomes composed of cholesterol (Avanti Polar Lipids), 1,2-dioleoyl-sn-glycero-3-phosphocholine (DOPC; Avanti Polar Lipids), 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC; Avanti Polar Lipids), 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphoethanolamine (POPE; Avanti Polar Lipids), disialoganglioside GD1a from bovine brain (Sigma-Aldrich), and N-(fluorescein-5-thiocarbonyl)-1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine, triethylammonium salt (Fluo-4Invitrogen) in a molar ratio of 20:20:20:40:1.64 × 10^-3 were prepared as follows. Purified lipids dissolved in chloroform, with the exception of GD1a, which was dissolved in a 1:1 chloroform:methanol mixture, were combined in the molar ratio listed above and dried to a film under an argon gas stream. The film was further dried under vacuum for 2 h. The film was then resuspended in HNE-50 pH 7.4 at 20 mg/mL by five freeze-thaw cycles in liquid nitrogen. The resuspended lipid solution was extruded 21 times at 40 °C through a polycarbonate membrane with a pore size of 0.2 μm (Whatman) to form liposomes. The liposomes were diluted fourfold in HNE-50 pH 7.4 and flowed into a flow-cell channel. The liposomes were incubated in the flow cell at RT for 10 to 30 min, during which time they adsorbed to the glass, fused with neighboring liposomes, and ruptured to form a supported lipid bilayer on the glass coverslip.

**Imaging Single-Particle Hemifusion.** Labeled virus was diluted 50-fold in HNE-50 pH 8.0 or HNE-50 pH 7.4 and flowed at a rate of 0.04 mL/min into a flow cell containing a supported lipid bilayer until the desired density of virus on the bilayer was achieved. For experiments with an initial pH value of 6.6, labeled virus in HNE-50 pH 7.4 was flowed into the flow cell and then washed and incubated in HNE-50 pH 6.6 for 5 to 10 min. To initiate hemifusion, low-pH buffer was continuously flowed into the flow cell at a rate of 0.06 mL/min. The flow-cell channel was illuminated in TIRF mode simultaneously with a 488-nm solid-state laser (Coherent) and a 561-nm solid-state laser (Coherent) through an oil-immersion, high-numerical-aperture objective (N.A. 1.45). Fluorescence emission was collected through the same objective, filtered through a dual-band-pass filter (Chroma Technology), and recorded by an EMCCD camera (Hamamatsu ImageEM) at a frame rate of 5 Hz for 250 to 300 s. Laser powers of 30 W for the 488-nm line and 5 W for the 561-nm line, as measured on the laser table, were used. Transmittance through the objective was 53% for the 488-nm line and 56% for the 561-nm line. All experiments were conducted at room temperature.

**Imaging Single-Particle Rolling.** Labeled virus was diluted 50-fold in HNE-50 pH 8.0 and flowed at a rate of 0.04 mL/min into a flow cell containing a supported lipid bilayer until the desired density of virus on the bilayer was achieved. At a flow rate of 5 Hz for 250 to 300 s, laser powers of 30 W for the 488-nm line and 5 W for the 561-nm line, as measured on the laser table, were used. Transmittance through the objective was 31% for the 488-nm line and 34% for the 561-nm line. All experiments were conducted at room temperature.

**Data Analysis.** The single-particle hemifusion data were analyzed as previously described (6). To analyze the single-particle rolling data, images were sharpened by convolution with a Mexican hat filter and smoothed by a median filter (pixel size of 2) using ImageJ software (NIH). Particle locations were picked manually in the first frame of the recorded movie. A rectangular region of interest (ROI) was defined for each particle in each frame of the movie, extending in the direction of flow from the initial particle location in the first frame to the edge of the field of view. These ROIs were used to construct position-versus-time kymographs for each particle, and the particle track was traced manually. Particle velocities were determined from the slopes of the traced particle tracks. The kymograph and velocity analyses were performed using custom-written software in MATLAB (MathWorks).

**Negative-Stain Electron Microscopy.** VSV particles were incubated in buffers HNE-50 pH 7.6, HNE-50 pH 7.0, and MES-50 pH 6.6 for at least 15 min at room temperature at a particle concentration of 0.05 to 0.1 mg/mL. The samples were adsorbed to carbon-coated collodion-support grids for 30 s, blotted, rinsed once in 2% (wt/vol) phosphotungstic acid (PTA), blotted, stained for 15 to 30 s with 2% (wt/vol) PTA, blotted again, and dried under light vacuum. The grids were glow-discharged before sample adsorption. The pH of the phosphotungstic acid was adjusted with sodium hydroxide to match the pH of the incubating buffer. Samples were examined using a JEOL 1200EX electron microscope operated at 80 kV (Department of Cell Biology Electron Microscopy).
Facility, Harvard Medical School). The mean thickness of the G-protein layer and the SD are reported in Results.

**Gₐ Purification.** Gₐ was cleaved and purified from virus particles as described in ref. 21 with the following modifications. In the cleavage reaction, the concentration of WT virus was 10 mg/mL of G-W72A virus 11.1 mg/mL, and of thermolysin 0.6 mg/mL. Virus concentration was measured by Bradford assay (Bio-Rad) using a BSA standard curve. Total reaction volumes ranged from 300 to 700 µL. Proteolysis was stopped by the addition of both blocking buffer (900 mM Tris-HCl, pH 8.8, 50 mM EDTA) and protease inhibitor mixture (Complete, EDTA-free; Roche). Cleavage reactions were then spun through 20% (wt/vol) sucrose cushions (20% (wt/vol) sucrose, 20 mM Tris HCl, pH 8.8, 10 mM EDTA) in a TLS-55 rotor (Beckman Coulter) at 48,000 rpm for 1 h at 4 °C. Supernatants were diluted 1:10 in buffer A (10 mM Tris-HCl, pH 8.8, 10 mM EDTA) and loaded onto an anion-exchange column (HiTrap Q HP 5-mL column; GE Healthcare). Gₐ was eluted with a linear gradient of buffer B (10 mM Tris-HCl, pH 8.8, 1 M NaCl, 10 mM EDTA) at μₑₐ₄ from −21 to 23% buffer B. Purified Gₐ was concentrated through an Amicon Ultra-4 centrifugal filter unit with a 10-kDa molecular-weight cutoff (Millipore). Final Gₐ concentration was determined by densitometry of Coomassie-stained gels using a BSA standard curve. Purified Gₐ was stored at 4 °C and used within 3 to 4 d for liposome-fusion experiments.

**Liposome Association of Gₐ.** The liposome-association assay was modified from a previously described protocol (31). Liposomes were prepared as described in the previous section but resuspended at a concentration of 10 mg/mL in HNE-50 pH 8.0, HNE-50 pH 7.6, MES-100 pH 6.6, and MES-100 pH 6.0 buffers. All solutions used in the liposome-association assay, including the liposomes and the cross-gradient solutions, were supplemented with EDTA such that the final EDTA concentration was 5 mM. Five microliters of either Gₐ/WT (0.53 µg/µL) or Gₐ/W72A (0.52 µg/µL) was mixed with 20 µL liposomes and 125 µL of the matching pH buffer for a total reaction volume of 150 µL. The reactions were incubated on a Labquake at 1 h at 37 °C. The reactions were then mixed thoroughly with 200 µL of 70% (wt/vol) sucrose solution to yield a final sucrose concentration of 40% (wt/vol). To form a discontinuous sucrose gradient, the association reaction in 40% (wt/vol) sucrose solution was layered at the bottom of a centrifuge tube. Nine hundred microliters of 25% (wt/vol) sucrose solution was layered on top and 150 µL of 5% (wt/vol) sucrose solution was layered on top of that. Each sucrose solution was made in the pH buffer matching that of the association reaction (i.e., HNE-50 pH 8.0, HNE-50 pH 7.6, MES-100 pH 6.6, or MES-100 pH 6.0 buffer). The gradients were spun in a TLS-55 rotor at 52,000 rpm for 2.5 h at 4 °C. Two-hundred-microliter fractions were collected from the top of the gradient using wide-bore pipette tips. The fractions were stored at 4 °C before SDS-PAGE and immunoblotting.

In the case of the reversible-reaction samples, the initial association was done in a reaction volume of 100 µL (5 µg Gₐ/WT = 20 µL liposomes + 75 µL matching pH buffer). After the 1-h incubation at 37 °C, 50 µL of 1 M Hepes (pH 7.0) was added to each reaction to give a pH of 7.0. Subsequent procedures were identical to those given above for the irreversible reactions. To trimerize, three G* monomers must be adjacent to each other on the surface of the virus. Therefore, the concentration of prep³ depends both on the concentration of G* and on the probability that the G* molecules are in the correct geometry. We used a Monte Carlo simulation (implemented in MATLAB) to determine the probability that G* would be in a geometry allowing trimerization. We constructed patches containing m monomers of G (where m equals three times the patch size P on the surface of the virion), seeded the patch with different numbers of G* monomers, ranging from 0 to m, and counted the number of trimers on each patch. We assumed that each G* monomer is potentially a member of only one trimer. We simulated 500 patches for each seed value of G* and calculated the probability that G* could trimerize for that seed value of G*. We fit a Gaussian function to our simulated probabilities to obtain the probability distribution for prep³ given the concentration of G*. We found that the probability distributions converged for patch sizes greater than 30 monomers. Eq. 3.2 is a Gaussian fit to the converged-distribution for the concentration of prep³ (Fig. 53). The final and third reaction is irreversible hemifusion requiring a concerted conformational change of a cluster of n extended G* trimers within a patch on the surface of the virion (Fig. S3A and reaction 1.3).

Based on these reactions, the kinetic differential equations describing the time-dependent changes of each of these G species are also shown in Table 1 (Eqs. 2.1 to 2.4). The pH in the system was modeled as a sigmoidal curve, where the pH dropped from pHŒ–WT = 7.0 to pHŒ–WT = 6.0 over a period of about 3 min (Eqs. 3.1, 3.2, and 3.3). We chose a sigmoidal curve over a step function for better numerical integration of the differential equations (see below). Eq. 3.4 relates pH and [H⁺]. Whereas [G*], [H⁺], [G*], [prep³], and [G*] are local concentrations of the species on the virion surface in mol-1 L⁻¹, T is the number of G* trimers in a patch with size P, and Vmax and Vt is the number of total and fused virions, respectively. We used Eq. 3.5 to calculate T from [G*], where P is the maximal number of trimers for a given patch size. In a population of virions, however, different virions may be able to form different numbers of G* clusters, and virions with more G* clusters will have a higher probability of hemifusing than virions with fewer G* clusters. Therefore, over a population of virions, the rate of hemifusion is the weighted sum of the individual fusion rates of virions containing different numbers of clusters, from 0 to Cmax where the weights are the probabilities that a virion contains i clusters at a given number T, which is P(r=i)cclusters (i) T in Eq. 2.4. To find these probabilities, we again used Monte Carlo simulations (implemented in MATLAB). We simulated the virion as a roughly circular patch of P points on a hexagonal array for P = 13, 31, 55, 73, and 109. Each point represented the position of a potential G* trimer on the virion surface. For a given [G*], we calculated the number of G* trimers that would correspond to on the patch and simulated 10,000 virions in which the number of points on the patch was randomly assigned a G* conformation. Clusters of adjacent G* trimers were then detected on each simulated virion. We defined clusters of three, four, five, and six adjacent G* trimers (n = 3, 4, 5, or 6) as previously outlined in ref. 7. The numbers of simulated virions containing 0 to Cmax clusters were tallied separately and divided by the total number of virions to obtain the probabilities. For computational efficiency, the probabilities were calculated for all possible T for a given patch size, where each column was the probability of hemifusing for patch size P and each row was the probability of forming i clusters at that concentration of preG. We numerically integrated the differential equations using the odeint function implemented in the SciPy Python library (version 0.17.0) (https://www.scipy.org/). The goal of model optimization is to find the set of parameter values that gives the best fit of the model to the observed data. We measured the quality of fit by calculating the root-mean-square deviation (rmsd) between the numbers of fused viruses at each time point in the model and experiment. We globally matched all nine regimes of initial and final pH simultaneously. Finding the global minimum of the multidimensional model function is nontrivial, and we therefore used a two-step approach. First, we ran simulations for a grid of parameter values, which explored five or six values for each parameter in the following range: prep = 6.5 to 7.3, K₁ = 100 to 800,000 mol⁻¹ L⁻¹ s⁻¹, k₂ = 0.01 to 10 s⁻¹, k₃ = 0.01 to 10 s⁻¹; patch sizes of P = 13, 31, 55, 73, and 109; and cluster sizes of n = 3, 4, 5, and 6. In a second step, we further optimized the best fit of the initial grid search for each combination of P and n using the SciPy minimization function with the Nelder-Mead algorithm (32).

ACKNOWLEDGMENTS. We thank Anna Loveland and Joe Loparo for help with TIRF microscopy and discussions; Amy Lee and David Cureton for help with virus growth and purification; Erick Matsen and Kyung-Suk Kim for initial discussions on the model; Maria Ericsson and the Harvard Medical School Cell Biology Electron Microscopy Facility; and Raffaele Potami for help with setting up parallel computing. I.S.K. acknowledges a National Science Foundation Graduate Research Fellowship. The research was supported by NIH Grant CA-13202 to S.C.H., who is an Investigator in the Howard Hughes Medical Institute.
Kim et al.