Characterization of the water defect at the HIV-1 gp41 membrane spanning domain in bilayers with and without cholesterol using molecular simulations

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1. Introduction

One of the proteins responsible for mediating fusion of human immunodeficiency virus 1 (HIV-1) particles to target cells is known as gp41. It resides in the viral membrane in so-called envelope spikes, which are trimers of noncovalently-attached dimers of envelope glycoproteins gp120 and gp41 [1,2]. In the native spike, gp41 spans the viral membrane as the spike’s anchor and is thought to be hidden in a metastable conformation by gp120. The membrane spanning domain (MSD) of gp41 is of great interest as it is responsible for both anchoring gp41 in a stable conformation by gp120 and for playing a role in fusion, according to truncation studies [3,4]. Mutagenesis has also shown that the specific sequence of the twenty-seven residue MSD is important for membrane fusion [5–11]. The MSD houses the GXXXG motif, postulated to mediate transmembrane helical interactions, and the charged residues K681, R694, R705, and R707 (Table 1) [3–14].

The gp41 MSD is predicted to be membrane-spanning based on mutagenesis and to be α-helical based on both bioinformatic and CD analyses [10,15–17]. Recently, our group supported this conclusion using all-atom metadynamics simulations and further showed that the membrane-spanning, α-helical MSD’s midspan arginine (R694) likely “snorkels” in the inner leaflet by allowing the guanidino-group access to the layer of lipid headgroups [18]. The snorkeling configuration is not surprising partially because a charged residue in an ideally structured hydrophobic lipid bilayer core is highly energetically unfavorable [19]. Therefore, transmembrane arginines probably have mechanisms to compensate for the charge in the middle of a lipid bilayer, and those in the middle of membrane-spanning α-helices can nevertheless be solvated through a so-called “water-defect” [19–24]. Indeed, previous calculations of the free energy of insertion from solvent to lipid bilayer of a midspan-arginine-containing transmembrane peptide (not gp41 MSD) by molecular dynamics and experiments ultimately determined the energetic cost of insertion to be only a few kcal/mol with defect solvation [19,25,26].

Bilayer water defects are characterized by water molecules and often lipid headgroups that have encroached deep into the bilayer hydrophobic core in order to interact directly with polar/charged residues. Relatively large water defects have been studied by solid-state NMR and have also been observed in various simulation systems involving guanidinium ions, arginine analogs, and arginine containing transmembrane proteins (both β-barrels and α-helices) in various lipids [20,21,23,26–31]. It has not been previously investigated if HIV-1 gp41 MSD displays a water defect in the native spike complex, but an arginine at position 694 (HXBc2 sequence) is required for fusion [11,4]. Therefore, it stands to reason that if a water defect is present due to
R694, the defect may play a role in fusion as well. Additionally, the HIV-1 membrane is rich in cholesterol (up to 50 mol-%), which is well known to increase the thickness and to more order model bilayers in the liquid phase [32,33]. Cholesterol seems to have multiple roles during the HIV lifecycle. It is important for Gag assembly at the plasma membrane, co-receptor binding and of course, fusion [34–39]. Since cholesterol-depletion compromises fusion [36,40], it is of interest to investigate the influence of cholesterol on solvation by defect near HIV-1 gp41 MSD.

In this paper, as part of a segment-based approach to investigating structure/function relationships in gp41, molecular simulations are used to study particular aspects of R694-solvation of the α-helical gp41 MSD monomeric peptide in a model lipid bilayer. Our major finding is that solvation of R694 invariably requires penetration of water into the inner leaflet, establishing a classical water defect. This feature of R694 solvation is robust with respect to the peptide’s local lipid composition, including the presence of cholesterol, and it is initial-condition-independent. However, we find that cholesterol appears to modulate the behavior of the defect. For instance, the presence of cholesterol appears to focus bilayer thickness and water penetration fluctuations at the midspan-arginine-containing protein, whereas these fluctuations are not strongly correlated to protein position in the cholesterol-free bilayers. Based on these findings, we offer some speculation that a robust water defect near a gp41 trimer perhaps contributes to the metastability of the spike and viral membrane, bringing it closer to a stability limit that must be crossed for virus–cell fusion.

2. Methods

2.1. General

All-atom molecular dynamics (MD) simulations were performed with NAMD 2.8 [41] along with VMD 1.9 [42], the CHARMM force field [43,44] with recent lipid-based corrections [45,46], and explicit TIP3P water.

Preliminary systems were run for 100 ns with configuration parameters based on those recommended by the CHARMM-GUI Membrane Builder (grid spacing of 1 Å, nonbonded shifted from 10 to 12 Å, 2 fs timestep) [47]. These simulations were compared to the same systems run with more aggressive configuration parameters (grid spacing of 2 Å, nonbonded shifted from 8 to 9 Å, 1 fs timestep) to determine if we could decrease computational time without sacrificing accuracy. We were able to use these more aggressive parameters for the heterogeneous, cholesterol-containing simulations (WT1, etc., see below) but not for the homogeneous dipalmitoylphosphatidylcholine (DPPC) simulations (WT1ΔChol, etc., see below). The WT1 simulation run with less aggressive parameters (known as WT1c) is also analyzed for various observables in Supplemental Figures 1, 2E/F, 3A, 7, and 8, for comparison to the WT1 system. In all cases, the Langevin thermostat was set at a temperature of 310 K and the Langevin piston Nosé–Hoover barostat was set at 1 atm. The z-cell dimension was allowed to fluctuate independently of x and y, which were kept at a constant ratio. Periodic boundary conditions were employed. All simulations were run on TACC XSEDE resources.

2.2. Choice of lipids

Cholesterol-containing bilayers are used to represent a model of the HIV–1 viral membrane. A major lipidic component in the HIV liposome is sphingomyelin, a fully saturated lipid, and HIV envelopes contain up to 50% cholesterol [34]. Because no CHARMM parameters for sphingomyelin exist in the open literature, we chose to work with DPPC, also a fully saturated lipid. DPPC and SM undergo their main phase transition at nearly the same temperature [48,49]. At the physiologically relevant temperature of 310 K, 50/50 DPPC/cholesterol bilayers will exist in the liquid-ordered phase [50,51]. To the best of our knowledge, no one has measured the main phase transition temperature (Tm) of a pure DPPC simulation using the CHARMM36 lipid force field parameters. Using other force fields, Tm has been suggested to be between 300 and 320 K [52,53], which compares fairly well to the experimental temperatures of 314–315 K [54,55]. Our pure DPPC membranes are gel-like and order parameters show the DPPC bilayers to be slightly less ordered than the cholesterol-containing bilayers (see Supplemental Figure 3).

2.3. System setup

We generated initial states for the WT1, WT2, and WT3 systems using a steered-MD (SMD) approach to insert an α-helical gp41 MSD model in three distinct locations of an equilibrated lipid bilayer composed of 50% DPPC and of 50% cholesterol, and each system was subjected to several rounds of equilibration following the procedure detailed previously [18] before launching individual 300 ns NPT production runs. Because lipid diffusion is so slow on a 300 ns timescale, the lipids cannot rearrange enough for ergodic sampling of all possible lipid arrangements; hence local compositions around each protein are largely determined by the choice of the initial insertion point. Therefore, the three WT simulation systems differed mainly in the local bilayer composition near the protein (Supplemental Figure 4). The same SMD approach was used to insert an α-helical gp41 MSD model into an equilibrated lipid bilayer composed purely of DPPC. We refer to this system as WT1ΔChol, and it was subjected to 300 ns NPT production MD after equilibration. Two other replicas, WT2ΔChol and WT3ΔChol, were branched off from the WT1ΔChol trajectory (at 0 ns and pre-production). The protein backbone was restrained and the lipids were allowed to further equilibrate around the peptide for 25 ns before starting production runs. Both protein-free bilayers were initially generated using the CHARMM-GUI membrane-builder and equilibrated [47]. For comparison of some observables, the membranes (with no peptide) were also run for 300 ns NPT MD; these systems are referred to as DPPC and DPPC/Chol systems. The R694L mutant system was generated using the VMD mutator plug-in operating on the equilibrated WT1 system, and this mutant system was subjected to both metadynamics (detailed below) and 300 ns production MD. Post-production, R694L was mutated back to WT, generating the R694LR system, which was subjected to 300 ns NPT production MD. All systems were neutralized and brought to 0.1 M NaCl before production runs. The cholesterol-containing systems have ~58,000 atoms, including 183 cholesterol and 151 DPPC molecules. These systems are roughly 80 × 80 × 80 Å3. The pure DPPC systems have ~89,000 atoms, including 334 lipid molecules. These systems are roughly 90 × 90 × 100 Å3.

2.4. Metadynamics of R694L MSD

Following our previous work on WT MSD [18], we used metadynamics [56–59] to test whether or not the R694L MSD’s minimum free-energy state is α-helical in a 50% DPPC and 50% cholesterol bilayer membrane. Briefly, in metadynamics, the MD trajectory is biased by a history-dependent potential that is the sum of Gaussians deposited
along a chosen collective variable (CV); the forces arising from these Gaussians allow high-probability regions along CV space to be explored first and discourage the system from revisiting the same CV values. As before, we used as a CV the RMSD of the backbone (C, Cα, N, O) atoms of the peptide with respect to a perfect α-helix, an average over a total of 109 atoms. The lower boundary, an RMSD of 0 Å, indicates a perfect helix, and the upper boundary, an RMSD of 10 Å, indicates a partially unfolded peptide. The metadynamics parameters included a Gaussian width of 0.01 Å, a weight of 0.15 kcal/mol, and a deposition frequency of 1 ps.

2.5. Observables computed in the production molecular dynamics simulations

The mass density as a function of position along the global membrane normal direction, z, of various components [protein, cholesterol, lipid headgroups, lipid tails, water (local and global)] averaged over

![Fig. 1. Final cumulative, conformational potential of mean force (PMF) in kcal/mol vs. collective variable (RMSD of backbone compared to perfect helix) in Å from the R694L peptide metadynamics, in black. The PMF vs. CV from the WT1 peptide metadynamics from our previous study is also shown here, in red diamonds [18]. Error bars represent standard deviations of the last 20 ns.](image)

![Fig. 2. Collective variable (RMSD of backbone compared to perfect helix) in Å vs. simulation time in ns and histogram of RMSD vs. frequency for equilibrium MD of (A) WT1, (B) R694L, and (C) WT1ΔChol systems.](image)
the last 100 ns of the trajectory was determined after shifting the \(x,y\)-origin to the peptide center of mass, shifting the \(z\)-origin to the bilayer center of mass, and rewrapping all coordinates into the central image. Data for the WT MSD in the mixed bilayer was averaged over replicas WT1, WT2, and WT3, while the data for the WT MSD in the pure DPPC membrane was averaged over replicas WT1\(\Delta\)Chol, WT2\(\Delta\)Chol, and WT3\(\Delta\)Chol. The number of water molecules uniquely associated to each residue was determined by partitioning every water molecule within 4 Å of any protein atom into cells of a Voronoi tessellation defined by the locations of the \(\alpha\)-carbons (i.e., a water belongs to the residue to whose \(C_\alpha\) it is closest). Helix tilt was defined by the angle between the vector aligned along the helix and the global \(z\)-axis.

![Graphs showing mass density along membrane normal for various protein configurations](image)

**Fig. 3.** (A, C, and E) Mass density in \(10^2\) amu/Å\(^3\) along membrane normal in Å during equilibrium MD for various components of the system: protein, local water (defined as within 4 Å of protein), and midspan residue (arginine or leucine). (B, D, and F) Mass density in amu/Å\(^3\) along membrane normal in Å during equilibrium MD for various components of the system: lipid tails, lipid headgroups, water, and cholesterol. A and B are averages for WT1, WT2, and WT3 systems. E and F are averages for WT1\(\Delta\)Chol, WT2\(\Delta\)Chol, and WT3\(\Delta\)Chol systems. All statistics are from the last 100 ns of each trajectory and error bars in A, B, E and F represent standard error.
Tilt was averaged over 3 replicas for the WT systems in a mixed and pure DPPC bilayer and only included statistics for the last 100 ns. Maps of membrane thickness $L(x,y)$ were determined by dividing the $x,y$-plane into squares of size $4 \times 4 \text{Å}^2$, calculating the $z$-distance between the center of mass of the lipid headgroups in the outer and inner leaflets within each square and averaging over the trajectory. Also mapped in the plane of the membrane was the average-minimum distance sampled by any water molecule on the inner side to the global membrane midplane, at the same resolution as $L$.

In general for this paper, if an observable is averaged over the trajectory for a single system, we report the standard deviation. If an observable is averaged between replicas (ex. WT1, WT2, and WT3), we report the standard error. We label these as SD and SE, respectively.

3. Results

3.1. HIV-1 gp41 MSD and its R694L mutant span the cholesterol-containing DPPC bilayer as $\alpha$-helices

The result for the R694L MSD metadynamics is very similar to that seen for the WT1 MSD, [18] namely, the most stable state is an $\alpha$-helix. The primary output of the metadynamics calculation of the R694L MSD is the potential of mean force (PMF) as a function of RMS distance from an ideal $\alpha$-helix (chosen to be the collective variable, CV). The cumulative PMFs averaged from the last 20 ns for the WT1 (computed previously) [18] and R694L systems are shown in Fig. 1. Metadynamics required approximately 320 ns to achieve diffusive sampling of the CV and convergence of the PMF, as seen in Supplemental Figures 5 and 6. It is interesting to note that multiple unfolding–refolding events were observed.

The 300 ns production NPT MD simulations also support the conclusion that the $\alpha$-helical state is stable for both the WT1 and R694L systems. In Fig. 2A and B, we show traces of the metadynamics CV observed during equilibrium MD launched from helical initial states for both systems, along with CV histograms. Both systems remain close to perfectly $\alpha$-helical. The WT1 peptide seems to have greater flexibility than the R694L peptide, as indicated by its wider RMSD distribution. In Fig. 2C, we show similar data for the WT1ΔChol system, which indicates that the WT peptide is helical in a pure DPPC membrane.

Production MD further shows that these stable $\alpha$-helices remain in a classical membrane-spanning orientation. In Fig. 3, we show mass-density profiles, averaged over the last 100 ns, for various components in the WT (averaged over 3 replicas), R694L, and WT1ΔChol (averaged over 3 replicas) systems. The right-hand panels in this figure show global density profiles that establish the $z$-span (membrane normal) of the bilayer, and the accompanying left-hand panels show the scaled local density of protein and water within 4 Å of the protein. Clearly all proteins span the bilayer. Also highlighted in the left-hand panels of Fig. 3 are the mass density distribution of atoms in the 694 residue; we see that R694 density overlaps with water density within 4 Å, indicating the R694 is solvated in both the WT and WT1ΔChol systems. R694L is clearly not solvated, as expected.

3.2. The midspan arginine is solvated by a robust water defect

Fig. 4 shows end-point snapshots from the WT1, R694L, WT1ΔChol, and WT3ΔChol systems, which illustrate some of the major differences among these systems. The major difference between the WT1 and R694L configurations shown here is that the WT1 has a classical water defect (highlighted by cyan van der Waals spheres) that connects the R694 side-chain with the bulk solvent through the inner leaflet, while no such defect exists in the R694L system. Noticeable in the snapshots of the WT1ΔChol and WT3ΔChol configurations are the overall gel-like alignment of the DPPC tails except near the proteins, which also display classical water defects.

Fig. 4. Representative system configurations from equilibrium MD at 300 ns rendered in VMD [42] for the WT1, R694L, WT1ΔChol, and WT3ΔChol systems. Lipid headgroups are in red vDW, lipid tails in white vDW, cholesterol in yellow vDW, water in cyan isosurface, peptide in orange new cartoon, and R694 or R694L in orange vDW. Water with oxygens within 4 Å of the protein and located on the inner leaflet of the membrane are shown in cyan vDW. For clarity, lipids, cholesterol, and water in the foreground of all four configurations are not shown.
Solvation of R694 is a robust feature of the entire set of production MD runs on the WT MSD. In Fig. 5A, we show the number of water molecules uniquely associated with each residue of the MSD for the three wild-type systems in the mixed bilayer (WT1, WT2, and WT3, averaged over the final 100 ns of each simulation). The data show that the midspan arginine (local residue index 14) directly interacts with 4.8 ± 0.03 (SE) water molecules throughout the trajectory. Peaks every fourth residue thereafter signify the amphipathic nature of the sequence and the fact that the waters predominantly solvate one side of the helix and, importantly, connect to the bulk interior-side water. It is interesting to note that the uniformity in the solvation profile in the three systems exists in spite of large differences in the local membrane composition near the protein in the three systems, illustrated in Supplemental Figure 4.

Fig. 5B shows that the midspan leucine mutant has no water defect. Since the water defect in the WT1 system may have resulted from waters solvating R694 prior to the production run (see System Setup), a second WT system was generated by spontaneously “undoing” the mutation of R694L to generate the R694LR system which was allowed to equilibrate and accumulate 300 ns of production MD. Water molecules enter the membrane within 9 ns and the water defect re-establishes itself on a roughly 100 ns timescale, as shown in mass density plots in Supplemental Figure 2C, and D. In Fig. 5B, we show the number of water molecules uniquely associated with each residue of the MSD for the R694LR system, which is nearly identical to that of the original WT1 system. Because the R694L system has no water defect, the waters that form the defect in R694LR spontaneously entered the membrane upon mutation and production MD. As shown in Fig. 5C, the midspan arginine interacts with 6.7 ± 0.44 (SE) waters in the pure DPPC bilayer with an amphipathic solvation signature similar to those in the mixed bilayer.

Fig. 6A shows the tilt angle of the MSD helical axis with respect to the global z axis as a function of simulation time in the three mixed-bilayer WT simulations. Though strongly fluctuating, all MSDs in the mixed-bilayer sample around a tilt angle of 19.5° ± 1.4 (SE) over the last 100 ns. The data in Fig. 6B show that the midspan leucine mutant displays a lower tilt angle than the WT with an average of 14.3° ± 3.1 (SD) and the R694LR system has an average of 24.7° ± 3.8 (SD) for the last 100 ns. Taken together, this suggests that the water defect is correlated with larger tilt angles. In contrast to the uniformity of tilt angle observed for the cholesterol-containing systems, Fig. 6C illustrates that the tilt angles for the MSDs in the pure DPPC bilayers can vary significantly (averages of 7.2° ± 3.0, 22.6° ± 3.0, 39.5° ± 3.7 (SD) for WT1ΔChol, WT2ΔChol, WT3ΔChol, respectively). For example, WT1ΔChol shows a

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Fig. 5. Number of water molecules within 4 Å of protein vs. amino acid that each water molecule is uniquely attributed to during equilibrium MD. All statistics are from the last 100 ns of each trajectory and error bars represent standard deviation.

Fig. 6. Tilt angle in degrees vs. simulation time in ns.
nearly vertical MSD helix, a fact also illustrated by the snapshot of this system shown in Fig. 4. At the other extreme, WT3ΔChol samples tilt angles between 30 and 45°. Yet all three MSDs in pure DPPC show classical water defects, as indicated by the water counts in Fig. 5C. The apparent relationship between the water defect and helix tilt is therefore dependent on the membrane cholesterol content: only in the mixed bilayer is larger tilt unambiguously associated with the defect. To explain this, we next turn to local maps of membrane thickness.

3.3. Defect-mediated local thinning is cholesterol-dependent

The two left-hand columns of Fig. 7 show maps of membrane thickness $L$ (see Methods) and its standard deviation $\sigma_L$ averaged over each of the three contiguous 100 ns intervals of the 300 ns production runs for the mixed-bilayer system WT1 and the leucine mutant R694L. $L$ is nearly identical between the two mixed-bilayer systems, except for the area within about 10 Å from the protein in

![Maps of membrane thickness, L, (top) in Å and standard deviation, $\sigma_L$, (bottom) in Å for the WT1, R694L, WT1ΔChol, and WT3ΔChol systems during 100 ns intervals of equilibrium MD. Overlaid on the maps are the x and y positions of the non-hydrogen atoms of the peptide (black circles) and the 694 residue (white circles) from the last frame of the trajectories. The N- and C-termini are labeled.](image-url)
which \( L \) is significantly lower for the WT1 MSD. Fluctuations in \( L \) are essentially absent in the leucine-mutant system, while fluctuations of more than 7 Å appear correlated with the position of the water defect in the WT1 system. The time-course of these maps for both the mixed-bilayer systems is essentially constant, indicating that the uniformity of membrane thickness and the correlation of strong fluctuations with the water defect site are stable phenomena.

In the two right-hand columns of Fig. 7 we show \( L \) and \( \sigma_L \), in the same manner as in the left-hand columns, but for two of the pure-DPPC systems, WT1\( \Delta \)Chol and WT3\( \Delta \)Chol. \( L \) is much less uniform over the entire system for the pure DPPC bilayers, which display large regions of distinctly lower thickness associated with lipid tails that are not ordered. Bilayer thinning local to the peptide is also much less severe in WT1\( \Delta \)Chol than in the mixed-bilayer system WT1, which is consistent with the observation of near-zero tilt of the MSD helix in the pure DPPC system. However, we still observe that the strongest fluctuations in \( L \) are correlated with the water defect, but that these fluctuations in \( L \) require at least 100 ns to equilibrate. In contrast to WT1\( \Delta \)Chol, the MSD helix in WT3\( \Delta \)Chol is strongly tilted and lies in a locally thin region of the membrane. Nevertheless, fluctuations in membrane thickness are localized around the defect and these fluctuations also require at least 100 ns to equilibrate. Peristaltic motions, similar to our definition of thickness fluctuations here, have been measured experimentally and computationally in pure DPPC bilayers and the thickness fluctuation amplitude was found to be 3.5–5 Å (at various temperatures) [60–62]. These agree fairly well with our observations.

Because membrane thickness depends on locations of lipid headgroups in both leaflets, it does not necessarily reflect the true effect of water penetration into one bilayer. In Fig. 8 we show maps of the average-minimum distance of any water molecule from the inner side to the global membrane midplane (\( z = 0 \)), and its standard deviation, at 4 × 4 Å² resolution, for three representative systems (WT1, WT1\( \Delta \)Chol and WT3\( \Delta \)Chol). In the mixed-bilayer, water penetration is strongly localized at the protein and also strongly fluctuating. For the two pure DPPC cases, the areas of greatest water penetration are evidently unions of areas of lowest membrane thickness and the local water defect. That is, the midspan arginine is not solely responsible for bringing water close to the midplane; thickness fluctuations do so as well. It appears that the main effect of cholesterol is to localize thinning of the membrane and contain the water defect.

Maps of membrane thickness and average minimum water penetration for systems not shown in Figs. 7 and 8 are in Supplemental Figures 7 and 8. For example, in Supplemental Figure 7, the protein-free, cholesterol-containing bilayer, DPPC/Chol, has a similar uniform thickness of 48 Å compared to the protein-free areas of the water-defect-free R694L system. This thickness is consistent with previous experimental and computational studies on DPPC/Chol systems and is larger than cholesterol-free DPPC membranes in the liquid phase due to the ordering effect of cholesterol [32,33]. Also, the protein-free bilayer, DPPC, is consistent with previous studies on pure DPPC bilayers in the solid phase; this system has an area per lipid of 48.8 Å² comparable to the experimental value of 47.2 Å² [63].

4. Conclusion

We have used an extensive set of MD simulations to probe the nature of the water defect that solvates the midspan arginine (R694) in the HIV-1 gp41 MSD and its relationship to cholesterol. Our results support the previous finding that the thermodynamically preferred state of the excised gp41 MSD in a cholesterol-containing-bilayer is as a membrane-spanning \( \alpha \)-helix, and further metadynamics results indicate that the R694L mutant is also stable as a membrane-spanning \( \alpha \)-helix. The water defect solvating R694 is a robust feature of all simulations of wild-type gp41 MSD, independent of local membrane composition and initial conditions. Cholesterol's major effect seems to be localizing membrane thinning to the water defect, which appears to more tightly regulate the tilt angles displayed by membrane-spanning MSD helices. The defect-local membrane thinning conferred by cholesterol is strongly fluctuating.

The insights gained from these simulations allow us to embark on some informed speculation that we hope will stimulate further experimental work aimed at understanding structure–function relationships in spike-membrane systems. First, we speculate that tightly regulated tilt angles for the gp41 MSDs are important for stabilizing trimERIC complexes that involve direct interaction of MSDs, because an axisymmetric trimer with a three-fold axis normal to the bilayer plane would require its monomeric constituents (individual MSD helices) to have the same tilt angle. This may mean that cholesterol is in fact important in conferring stability to the HIV-1 envelope spike.

Second, the effect of the water defect on the stability of the membrane against poration or fusion could also be important. An interesting question to pursue now is, which represents a “weaker” membrane in the context of the HIV-1 gp41 MSD: the cholesterol-containing or cholesterol-free bilayer? Although we have not done the calculations to answer this question, it may be worth considering that cholesterol-depleted HIV-1 is unable to infect, and that this effect is reversible. It is possible that strong water fluctuations along membrane normal in

![Fig. 8](image-url)
the inner leaflet hydrophobic core near membrane spikes on HIV-1 does not appear as cholesterol is depleted, and can easily resume as cholesterol is resupplied. It is reasonable to think that with the gp41 trimer in native HIV-1 spikes, the monomer water defects have merged and represent a local metastability of the membrane that can be harnessed during fusion. We speculate that these localized water fluctuations are important for metastability and therefore are important for entry.

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Appendix A. Supplementary data

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References
