

Strain and rupture of HIV-1 capsids during uncoating

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Viral replication in HIV-1 relies on a fullerene-shaped capsid to transport genetic material deep into the nucleus of an infected cell. Capsid stability is linked to the presence of cofactors, including inositol hexakisphosphates (IP₆) that bind to pores found in the capsid. Using extensive all-atom molecular dynamics simulations of HIV-1 cores imaged from cryo-electron tomography (cryo-ET) in intact virions, which contain IP₆ and a ribonucleoprotein complex, we find markedly striated patterns of strain on capsid lattices. The presence of these cofactors also increases rigidity of the capsid. Conformational analysis of capsid proteins (CA) show CA accommodates strain by locally flexing away from structures resolved using X-ray crystallography and cryo-ET. Then, cryo-ET of HIV-1 cores undergoing endogenous reverse transcription demonstrates that lattice strain increases in the capsid prior to mechanical failure and that the capsid ruptures by crack propagation along regions of high strain. These results uncover HIV-1 capsid properties involved in their critical disassembly process.

HIV capsid | all-atom molecular dynamics | uncoating | strain

The mature capsids of HIV-1 are large fullerene-like protein complexes that are comprised of more than 1,000 copies of the capsid protein (CA) (1). During replication, viral particles that bud from host cells are initially immature and composed primarily of a spherical array of Gag proteins that link essential viral proteins and enzymes into a linear polypeptide chain (2). As the virus matures extracellularly, proteolytic cleavage of Gag releases CA, which self-assembles in a cone-shaped geometry and packages two copies of the retroviral genome and associated enzymes into the capsid. Mature capsids are deposited during the fusion of HIV particles with the plasma membrane, in which the lipid envelope and embedded proteins are lost, leaving behind the core of the HIV-1 virus, i.e., the capsid and contents, in the cytoplasm of cells.

Capsids play essential roles during replication by transporting viral genetic material deep into the host cell (3). Pores in the capsid can bind or import small molecules, including inositol phosphates and nucleotides (4, 5), and the binding of inositol hexakisphosphate (IP_6) increases the stable lifetime of the capsid and promotes the assembly of CA into fullerene structures (6, 7). Cryo-electron tomography (cryo-ET) and other techniques have recently demonstrated that viral cores are imported with apparently intact capsids across the nuclear pore of infected cells (8-11). Reverse transcription processes inside the capsid can rupture the core, as seen in both atomic-force microscopy and cryo-ET experiments (12, 13), owing to an increased internal pressure on the capsid during the conversion of RNA to DNA. Uncoating of the capsid is a critical replication event, releasing enzymes and nucleic acids that integrate a copy of the virus in the host genome. Yet little is known about the physical properties underlying capsid rupture.

To investigate the structural and mechanical properties of HIV-1 capsids that lead to rupture and disassembly, we performed large-scale all-atom (AA) molecular dynamics (MD) simulations of HIV-1 capsids, containing native cofactors including IP₆, and a ribonucleoprotein complex (RNP). Analysis of the strain induced on the capsid reveals spatially correlated patterns, indicating that the capsid collectively cracks open along regions of high strain rather than slowly disassembling. Local fluctuations in the capsid volume decrease concomitantly with increased strain, consistent with a mechanical rigidification of the capsid in response to IP_6 and the RNP. Calculated free energy landscapes also reveal shifts in the conformational ensembles of CA in response to increased strain. Cryo-ET imaging of in vitro reconstituted HIV-1 cores incubated with nucleotides then add further insight into the temporal sequence of events during disassembly to demonstrate that strain is maximal prior to the formation of cracks in the capsid. These results show that capsids are intrinsically strained and characterize the molecular processes by which viral cores rupture.

Results

Our AAMD simulations of HIV-1 cores contained a total simulation size ranging from 44 to 76 million atoms. In prior AAMD simulations (14, 15), a model for the empty capsid shell, enclosing only water with neither nucleic acid contents nor the binding of inositol phosphates (i.e., IP_6), was constructed from low-resolution cryo-ET. In contrast, we derived six atomic models from fullerene lattice maps of the complete

Significance

The mature capsids of HIV-1 are transiently stable complexes that self-assemble around the viral genome during maturation, and uncoat to release preintegration complexes that archive a double-stranded DNA copy of the virus in the host cell genome. However, a detailed view of how HIV cores rupture remains lacking. Here, we elucidate the physical properties involved in capsid rupture using a combination of large-scale all-atom molecular dynamics simulations and cryo-electron tomography. We find that intrinsic strain on the capsid forms highly correlated patterns along the capsid surface, along which cracks propagate. Capsid rigidity also increases with high strain. Our findings provide fundamental insight into viral capsid uncoating.

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The authors declare no competing interest.

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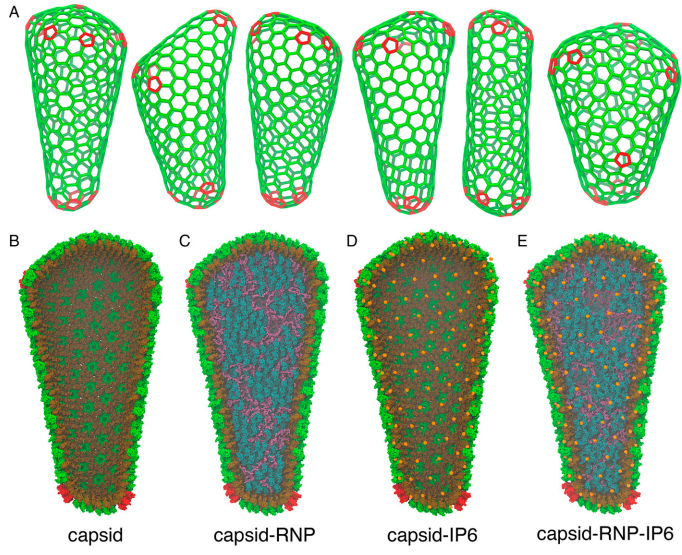


Fig. 1. Mature HIV-1 capsids are pleomorphic. (*A*) Fullerene geometries for the HIV-1 capsid were derived from cryo-ET images of intact virions (16). Atomic models for the capsid contain either (*B*) liquid water in the capsid interior, (*C*) a ribonucleoprotein (RNP) complex model, (*D*) IP₆ molecules bound to the capsid pores, or (*E*) both the RNP and IP₆. The CA NTD and CTD are colored in green and brown, whereas genomic RNA, nucleocapsid proteins, and IP₆ molecules are in purple, teal, and orange, respectively. Pentamer defects are colored in red.

capsid, with the positions and orientations of lattice components determined by an iterative alignment based on tomograms of intact HIV-1 virions (Fig. 1A) (16, 17). IP₆ polyanions were added to each CA hexamer or pentamer pore at the binding site, a location 2.7 Å above the R18 ring (5, 7). In the absence of atomic resolution into the structure of the RNP, we constructed a model of two copies of the 9-kb RNA genome in complex with nucleocapsid proteins consistent with experimental secondary structure constraints from selective 2'-hydroxyl acylation analyzed by primer extension (SHAPE) analysis (18) to mimic the HIV-1 RNP. Core particles were simulated at successively increasing levels of detail ranging from capsid shells containing only water to more realistic capsids containing the RNP model and IP₆ molecules (Fig. 1 B-E) (see Materials and Methods for a complete description). The C α positions of the atomic model for the capsids containing the RNP and IP₆ overlapped well with the 6.8-Å cryo-ET density of the CA hexamer and 8.8-Å cryo-ET density of the CA pentamer positioned in the lattice map (SI Appendix, Fig. S1). In aggregate, the AAMD simulations totaled 1.6 µs across the HIV-1 core particles (SI Appendix, Table S1).

Capsid Strain and Rigidity. During the AAMD simulations, the viral capsids remained intact. Empty capsids in bulk solvent without IP₆ did not dissociate or reassemble into more stable helical or spherical arrangements within the timescales simulated. Local deformations in materials and proteins have elucidated the mechanical properties of materials under stress and conformational changes resulting from protein-ligand interactions (19). To quantify these features, we computed per-particle strain tensors $(\boldsymbol{\varepsilon})$ for the centers of mass for each five amino acid residue segment in the CA domain. The volumetric strain $(s_{\rm V} = \frac{1}{3} {\rm Tr}(\boldsymbol{\epsilon}))$ measures the propensity for a particular region of the capsid to either swell or shrink (Fig. 2A). Empty cores containing liquid water had relatively little strain (CA; $\langle |s_V| \rangle =$ 2.4 $\times 10^{-2}$), which was distributed randomly across the capsid. Strain increased in the presence of the RNP complex (CA-RNP; $\langle |s_V| \rangle = 3.3 \times 10^{-2}$). IP₆ binding induced even larger effects that were also more spatially correlated than the capsid containing just the RNP (CA-IP₆; $\langle | s_V | \rangle = 3.6 \times 10^{-2}$). Cores containing an RNP complex and IP₆ molecules were the most strained, with patterns that formed unexpected striations (Fig. 24)

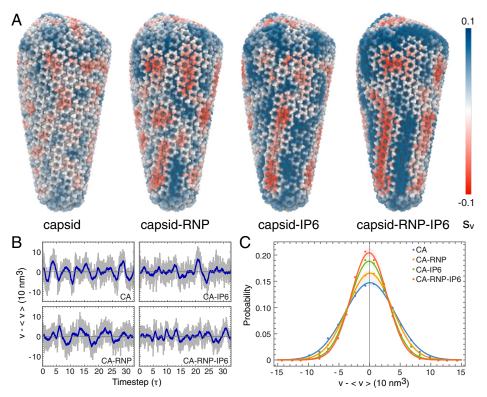


Fig. 2. Strain and fluctuation analysis of the HIV-1 capsid. (*A*) The first invariant of the strain tensor (s_V) or volumetric strain was computed for each capsid core complex. Red and blue colors correspond to compressive and expansive strain, respectively. (*B*) Fluctuations in the HIV-1 core volumes, measured as deviations from the average capsid volume ($V = 1.45 \times 10^5$ nm³). The subsampled timestep, τ , is 4 ns. Gray colors denote the instantaneous volume, whereas the blue line denotes a moving average within a one-timestep window. (*C*) Probability distributions for the volume fluctuation amplitudes in each capsid. Closed circles correspond to measured data, while the solid line indicates a Gaussian fit. The shaded regions denote the error in the distribution determined by block averaging.

along the capsid surface (CA-RNP-IP₆; $\langle | s_V | \rangle = 5.1 \times 10^{-2}$). Different capsid structures showed a similar trend of increased strain in the presence of IP₆ and RNP, but with slightly altered strain patterns that are attributable to differences between core morphologies (*SI Appendix*, Fig. S2).

To assess whether capsid properties change in response to IP₆ or RNP, we monitored the internal volumes of each core during the simulations. Although each pleomorphic capsid differed in size (Fig. 1A), average core volumes ranged from 1.15 to 1.53×10^5 nm³, with fluctuations of several tens of cubic nanometers about the equilibrium. Mean free volume fluctuations for the largest core are shown in Fig. 2B. Fluctuation amplitudes decreased markedly in the presence of either IP6 or the RNP, consistent with higher core rigidity (Fig. 2C, CA-IP₆ vs. CA-RNP) (SD: $\sigma_{CA} = 41.1 \text{ nm}^3$; $\sigma_{CA-RNP} = 36.3 \text{ nm}^3$; $\sigma_{CA-IP6} = 31.2 \text{ nm}^3$; $\sigma_{CA-RNP-IP6} = 29.2 \text{ nm}^3$). Decreases in the volume fluctuation amplitudes were consistent across cores of different morphologies (SI Appendix, Fig. S3). Fourier analysis showed a shift in the peak-to-peak frequencies of the dominant mode toward lower frequencies with the RNP, indicating that the capsids with RNP fluctuate more slowly (CA: v = 57.8MHz; CA-RNP: v = 39.4 MHz) (SI Appendix, Fig. S4). As the core rigidifies further upon IP₆ binding, the dominant lowfrequency mode broadens, and the fluctuations are distributed to higher-frequency modes, consistent with a stiffer capsid. Negatively charged IP₆ molecules bind tightly to an arginine ring in pores distributed throughout the capsid (4, 5, 7). RNP interactions with CA, on the other hand, could occur through interactions of positively charged residues on the flexible C-terminal domain (CTD) tail of CA with the RNP (7). The cofactor interactions at the pore and CTD tail change the

conformational flexibility of CA, resulting in an increase in capsid rigidity that introduces strain on the underlying lattice.

Conformational Analysis of CA Proteins. To examine the conformations CA proteins adopt in the actual capsid, we computed a three-dimensional (3D) free energy landscape (Fig. 3 A and B) from the spatial distribution occupied by the nonhydrogen atoms of CA across the pleomorphic lattice for the capsids containing IP₆ and RNP. At high contours ($\Delta G = 0.5$ kcal/mol), density for the protein backbone of CA is clearly visible, whereas at low contours ($\Delta G = 4.4 \text{ kcal/mol}$), variability in the ensemble of CA structures undergoing dynamical motion results in larger volumes (Fig. 3B). Less structured regions of the protein, including the CTD tail, β hairpin, and cyclophilin A (CypA) binding loop, had higher variability, and were more mobile. The hinge connecting the CA N-terminal domain (NTD) and CTD was more ordered than the other unstructured regions, possibly owing to NTD-NTD and CTD-CTD contacts in the adjacent CA domains of the capsid. The conformations of individual CA monomers in pentamers and hexamers resolved by cryo-ET (16) show small differences in the relative orientations of the NTD and CTD (Fig. 3C). Relative free energy differences between the conformations were quite small ($\Delta G < 0.2$ kcal/mol), indicating that CA can switch between the two states in the capsid under thermal motion.

Interfacial contacts between the NTD of one CA domain and the CTD of the adjacent CA domain differed appreciably in the pentamer and hexamer (Fig. 3 D-I). We employed an angle parameter, ϕ , defined as the angle between amino acid backbone centers of mass in the NTD helices and CTD helices, using the base of the NTD helices as a pivot, to assess the

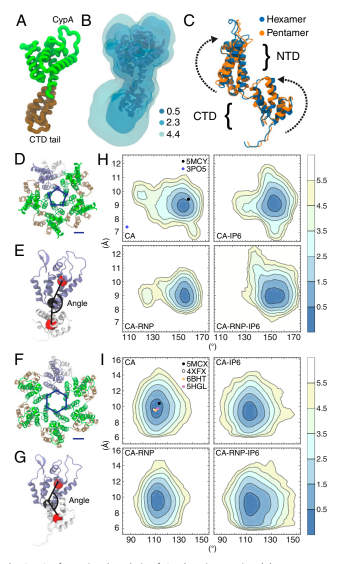


Fig. 3. Conformational analysis of CA domain proteins. (A) A monomer of the CA protein. (B) The 3D potential of mean force for CA conformations contoured at 0.5, 2.3, and 4.4 kcal/mol with respect to the Cartesian coordinates of the protein heavy atoms. (C) Differences in CA monomer conformations between the cryo-EM structure of the hexamer (PDB ID: 5MCX) and cryo-EM structure of the pentamer (PDB ID: 5MCY) consist of rotations in the NTD and CTD. The parameter, ξ , is used to describe the pore size and is defined as the center-of-mass distance between residues N21 and A22. The angle Φ is used to describe the relative orientation of a NTD and adjacent CTD and is defined as the angle between three centers of mass across the NTD and CTD (red and black spheres). ξ and Φ are shown for the CA hexamer (D and E) and pentamer (F and G). (H) (ξ , Φ) distributions for the hexamers in each capsid. (1) (ξ , Φ) distributions for the pentamers in each capsid. Contour lines correspond to increments of $f = -\log p$, where p is the probability. Closed circles correspond to (ξ, Φ) values from experimental structures in the Protein Data Bank.

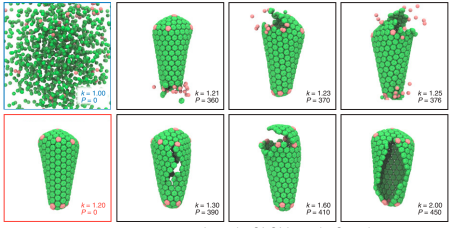
relative orientation of the NTD and adjacent CTD across hexamers and pentamers in the capsid during MD (Fig. 3 *E* and *G*). The angle, ϕ , was larger in the cryo-ET structure (16) of the pentamer ($\phi = 157^{\circ}$) than in the hexamer ($\phi = 113^{\circ}$), reflecting a greater curvature of pentamers over hexamers. Pore sizes were also monitored using a parameter, ξ , defined as the distance between the centers of mass of N21 and A22 on adjacent CA monomers (Fig. 3 *D* and *F*). Pentamers in flexible, empty capsids had high curvatures and small pore sizes (ξ_{CA}, ϕ_{CA}) = (7–11Å, 115–167°), although a subpopulation

(f = 2.5 - 3.5) shifted to a flatter curvature closer to that of the X-ray crystal structure of a pentamer (20) with engineered disulfide cross-links between N21 and A22, Protein Data Bank (PDB) identification code (ID): 3PO5 $(\xi, \phi) = (7.5\text{\AA}, 107^\circ)$ (Fig. 3H). IP₆ binding led to a conformational shift in pentamer distributions toward larger pore sizes ($\xi_{\text{RNP-IP6}} = 7-12.5\text{\AA}$), and capsids with both IP₆ binding and the RNP increased pentamer curvature ($\phi_{\text{RNP}} = 119 - 168^\circ$, $\phi_{\text{IP6}} = 121 - 170^\circ$, $\phi_{\text{RNP-IP6}} = 128 - 171^\circ$). CA hexamers had wider pores but flatter curvatures (lower ϕ) (ξ_{CA} , ϕ_{CA}) = (6.5–15.6Å, 92–136°) compared to pentamers. IP₆ binding and the RNP did not significantly affect pore size in hexamers, although there was a greater variation in hexamer curvature ($\phi_{IP6} = 90 - 147^\circ$, $\phi_{\text{RNP-IP6}} = 91 - 150^{\circ}$) owing to the strain induced on the overall capsid lattice (Fig. 31). X-ray crystallographic and cryoelectron microscopy (cryo-EM) structures agree with the minima of both curvature and pore size distributions, and resided within the lowest two contours of the (ξ, ϕ) distributions (f < 1.5), but CA conformations deviate away from these minima under increasing lattice strain.

The NTD-CTD interface is the binding site for phenylalanine glycine motifs required for nuclear import of the capsid (21) and small-molecule inhibitors including PF74 (22) and Lenacapavir (23) that disrupt HIV function. Our simulations indicate these binding pockets are modulated by the presence of IP₆ and the RNP. CA hexamers are less perturbed than pentamers but exhibited conformational shifts toward increased ϕ angles and larger binding pockets. Pentamers, on the other hand, existed in two dynamic subpopulations—one of which was consistent with the X-ray structure of a pentamer, and another consistent with a curved pentamer found in intact virions. The presence of strain (e.g., by IP₆ and RNP) induced local changes in the CA lattice that favored highly curved pentamers, and a more open binding site at the NTD-CTD interface.

Mechanical Rupture of the HIV-1 Capsid. Coarse-grained (CG) simulations in which the internal pressure on the capsid was varied were performed to examine how CA-CA interaction strengths altered capsid disassembly behavior (Fig. 4). To uniformly increase the pressure inside the capsid, particles with excluded volume interactions were added in the capsid interior until the point of rupture. Below a critical degree of CA-CA interaction (k = 1.20 kcal/mol), the capsid disassembles even without any internal pressure (P = 0), whereas above this interaction strength, the capsid remains stable, indicating that there is a minimum lattice energy required to stabilize the fullerene geometry of the CA lattice. The pressure required for rupture increased with increasing CA-CA interaction strength. At low CA–CA interaction strengths (k < 1.30 kcal/mol), small separations in the lattice lead to the gradual dissociation of CA and disassembly of the capsid along the separations. In the CG simulations, pentamers dissociated more quickly from the lattice, owing to the fewer attractive contacts the pentamers had in the lattice compared to hexamers. At higher CA-CA interaction strengths ($k \ge 1.30$ kcal/mol), rupture of the capsid was more collective, with larger cracks that formed across the capsid surface. These CG simulations highlight that the intermediates formed during rupture of the capsids are sensitive to the strength or rigidity of CA-CA interactions in the capsid.

Core Rupture during Endogenous Reverse Transcription. Cryo-ET imaging and lattice mapping were used to probe how core structure changed during reverse transcription (13). In brief, HIV-1 cores were released from purified virions by permeabilizing the viral membranes with the pore-forming melittin peptide and then stabilized by the addition of IP₆ at native cellular concentrations. Reverse transcription was initiated by adding



Increasing CA-CA Interaction Strength —

Fig. 4. Rupture of the capsid in CG simulations. A series of snapshots shows the mechanical failure of the capsid as internal pressure is increased until the point of rupture, taken from the trajectories at time $\tau = 10^6$ CG MD timesteps. The strength of CA–CA interactions (*k*) is given in units of kcal/mol, whereas the pressure (*P*) is given in units of spherical particles that are added to the capsid interior. At *k* = 1.00 kcal/mol; *P* = 0, the capsid slowly disassembles (blue panel). At *k* = 1.20; *P* = 0, the fullerene lattice remains stable (red panel). With low CA–CA interaction strengths (*k* = 1.21, 1.23, and 1.25 kcal/mol), small lattice separations lead to a gradual disassembly of the capsid. With high CA–CA interaction strengths (*k* = 1.30, 1.60, and 2.00 kcal/mol), rupture is characterized by the formation of larger cracks in the capsid.

deoxynucleotide triposphates (dNTPs) at concentrations found in CD4⁺ T cell cytoplasms (24). After 4 h of incubation at 37 °C, a total of 14 cores were found in the tomograms and mapped. Capsid structures showed a range of intermediates from intact cones to partially cracked capsids to nearly completely disassembled (Fig. 5 A, 1–14) capsids. We examined lattice separations in each core using a local order parameter, γ , which measures the near-neighbor contacts of a particle. In our AAMD simulations, lattice separations correlated with the strain, as expansive strain caused slight but measurable deviations in lattice separation ($\chi < 1.0$), whereas compressive strain induced more closely packed lattices ($\chi > 1.0$) (Fig. 5B). No neighboring particles are present at low values of χ , which provided a quantitative metric for the degree of cracking across the core ($\chi < 0.55$). Interestingly, tubular pill-shaped capsids (Fig. 5 A, 1) were nearly perfect fullerenes with no cracks present, whereas other, more conical capsids had cracks that were not visually obvious prior to computing χ in initial inspections (Fig. 5C). The various defects found in the experimental capsid structures included small ruptures along the narrow and broad ends of the fullerene cone as well as large cracks that formed along the length of the capsid (SI Appendix, Fig. S5). Intermediate values for χ correspond to small deviations and separations that reflect the expansive strain on the capsid $(0.75 < \chi < 0.9)$. Analysis of χ at these intermediate values revealed a bimodal distribution (Fig. 5D). We observed largely intact capsids with minimal strain (Fig. 5 A, 1 and 2) and high strain (Fig. 5 A, 3 and 4). Small ruptures or cracks in the lattice decrease the strain (Fig. 5 A, 5–8), which then increases during further stages of disassembly (Fig. 5 A, 9-11). We interpret these results to indicate that mechanical failure of the capsid and loss of capsid integrity lowers the strain, which then increases as the preintegration complex exits the core, and that reverse-transcribing capsids are strained prior to and during rupture of the capsid.

Discussion

We simulated entire HIV-1 core particles derived from cryo-ET at an atomic level of detail and computed the intrinsic strain on the capsid induced by IP_6 and RNP. Unexpected and correlated strain patterns formed on the capsid surface, and analysis of core volume fluctuations showed that the capsid mechanically

rigidifies in response to the presence of these cofactors. Our conformational analysis of CA proteins in the capsid also indicates that pentamers and hexamers have a surprising degree of dynamic flexibility that changes under strain. The CA pentamer adopts two distinct states, one corresponding to a state with a flat curvature and another corresponding to a curved state with a shift toward more highly curved conformations under increased strain. CG MD of the capsid under pressure demonstrates that rupture is initiated by small separations in the CA lattice that lead to disassembly of the capsid at regions of high strain, and as the CA-CA interaction strength is increased in the model, the rupturing becomes more collective, with large cracks forming in the capsid. Cryo-ET of core rupture during endogenous reverse transcription reveals that both small separations and large cracks in the capsid are present, and shows that lattice strain is locally maximal prior to mechanical failure. The capsids rupture in a fashion consistent with the propagation of cracks along highly strained regions, as observed in the simulations.

In biochemical assays (13), overstabilization at high IP_6 concentrations results in cores that do not produce reverse transcription products, and the binding of small-molecule inhibitors such as Lenacapavir (23, 25) can weaken capsid integrity and accelerate fracturing of the capsid (13), suggesting HIV-1 cores have physical properties that can be altered to disrupt retroviral life cycle processes. During trafficking from the cytoplasm to the nucleus of infected cells, viral capsids are exposed to interactions with a variety of host proteins. CypA binds to and enhances stability of the capsid in a concentration-dependent fashion (26) in the cytoplasm. Nucleoporins bind to and import viral cores across the nuclear pore of host cells (27, 28). Cleavage and polyadenylation specific factor 6 interacts with capsid complexes (22) and is involved in nuclear import and trafficking to integration sites (29, 30). Could these interactions with host proteins along with increasing internal pressure from reverse transcription weaken capsid integrity and rupture the capsid? Additional investigations should therefore further probe the molecular mechanisms involved in viral core interactions within the host cell.

Materials and Methods

AA Models of the Capsid Core. Initial atomic models for the CA hexamer and pentamer were constructed from the cryo-ET structure of the CA hexamer (PDB ID: 5MCX) and CA pentamer (PDB ID: 5MCY) derived from intact virus

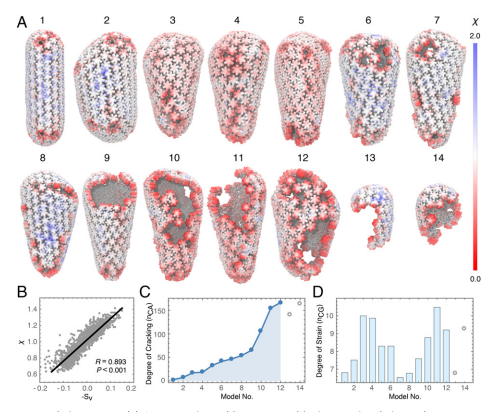


Fig. 5. Cryo-ET of HIV-1 cores during rupture. (A) Cores were imaged by cryo-ET and lattice mapping during endogenous reverse transcription. Each structure (1 to 14) is colored by a local order parameter (χ) that quantifies the degree of separation in the lattice. An ideally spaced particle in the lattice has a separation of $\chi = 1.0$. Red colors correspond to particles further apart on average from near-neighbors, whereas blue colors correspond to particles closer to near-neighbors. (*B*) Correlation between the strain (s_V) and lattice separation (χ) measured in the MD simulations. (*C*) The degree of cracking in each capsid model measured by the number of CA monomers containing a particle with $\chi < 0.55$. (*D*) The degree of strain in each capsid model measured by the number of particles (10³) with 0.75 < $\chi < 0.9$. Capsids 1 to 14 in A were ordered from no fractures to completely ruptured. Outliers in which the capsid was mostly disassembled are shown in open circles.

particles. Amino acid side-chain conformations were modeled based on the X-ray crystal structure of the CA hexamer (PDB ID: 4XFX) and disulfide-crosslinked CA pentamer (PDB ID: 3P05). Missing protein backbones residues were built using MODELER (31), and missing side chains were built using SCWRL4 (32). CTD tails for CA were transplanted onto the model using the NMR structure for the tail (residues 220 to 231) (PDB ID: 2KOD) onto the model. Atomic models for the CA hexamer and pentamer were then CG at a resolution of one amino acid residue per CG particle. Each CG hexamer and pentamer subunit was constrained as a rigid body, and positioned at the Cartesian coordinates and Euler angles that maximized overlap between the atomic model and the cryo-ET lattice map derived from intact virions (16).

CG models of the capsids were briefly relaxed in a 20-ps Langevin dynamics run under the canonical (NVT) ensemble with the large-scale atomic/molecular massively parallel simulator (LAMMPS) (33). To maintain overall protein shape, each CG particle included excluded volume interactions using a soft cosine potential $U_{Vol} = A [1 + \cos (\pi r/r_c)]$, when $r < r_c$. The cutoff (r_c) is the onset of excluded volume repulsion between CG particles, and was set to the separation distances found in the CTD interfaces if the distances were less than 10 Å, or otherwise 10 Å. The parameter A was taken to be A = 100 kcal/mol. The mass of the protein was evenly distributed among the CG beads. CTD contacts were maintained using an elastic network model that connected the two groups of CG particles in neighboring CTD dimers (residues 178 to 194) in a cutoff distance, $r_{\rm cut}$, with harmonic bonds; the potential energy of each bond is $U_{\text{bond}} = K_{\text{bond}} (r - r_0)^2$, where r is the separation distance and r_0 is the equilibrium bond length, set to the distance found in the crystal structure ($r_{cut} = 10$ Å, $K_{\text{bond}} = 0.1 \text{ kcal mol}^{-1} \text{ A}^{-2}$). Temperature was maintained at 300 K with a Langevin thermostat with a damping constant ($t_{damp} = 5$ ps). AA models for each CA hexamer and pentamer were then aligned with each CG capsomere subunit to construct an initial AA model for the capsid. Six complete capsids were constructed corresponding to the fullerene geometries derived from the cryo-ET structures two of which were selected for additional modeling.

For each of the capsid systems selected, IP_6 molecules were placed at positions corresponding to the bound X-ray crystal structure, 3.5 Å above an R18

ring that lines the pore of the CA hexamer and pentamer. Approximately 200 IP₆ molecules were used for each system, corresponding to a single IP₆ molecule for each capsomere subunit in the fullerene structure. A model of the ribonucleoprotein core was generated using a simplified CG model of RNA. The fulllength HIV-1 genomic RNA sequence was used (18), and CG particles were added to the capsid interior at a resolution of four beads per nucleotide base pair. The CG particles were labeled corresponding to the atom types in each base pair (adenine: C3' C8 N6 C2; cytosine: C3' C6 O2 N4; guanosine: C3' C8 O6 N2; uracil: C3' C6 O2 O4) to generate the viral RNA. Bonded interactions were added using harmonic restraints between the neighboring nucleotides to $(K_{\text{bond}} = 1.0 \text{ kcal mol}^{-1} \text{ Å}^{-2})$ and set to the average equilibrium distances between CG particles found between the represented atom types in the X-ray crystallographic structure of a model RNA template (PDB ID: 4GXY) to maintain geometric shape. Secondary structure constraints and base pairing interactions were implemented using harmonic restraints between the nearest-neighbor nonbonded CG particles between base pairs on the basis of high-throughput SHAPE reactivity data (18) ($K_{ss} = 0.1 \text{ kcal mol}^{-1} \text{ Å}^{-2}$). Two copies of the 9-kb genome were modeled and positioned in the interior of the capsid. CG particles for the nucleocapsid proteins were mixed with the RNA at random positions in the capsid interior, until there was a 1:1 correspondence between the number of CA and nucleocapsid for the RNP. The CG RNP model was energy minimized and relaxed under a brief 100-ps Langevin dynamics run. AA models for the RNA and nucleocapsid proteins (PDB ID: 1A1T) were then fit to the CG RNP, and subsequently energy-minimized and equilibrated as described below.

AAMD Simulations of Viral Capsid Cores. Solvated capsids were of sizes ranging from 44 to 76 million atoms including water molecules and ions. Na⁺ and Cl⁻ ions were added to the bulk solution until the salt concentration was 150 mM NaCl to produce an electrostatically neutral system. Periodic boundary conditions were imposed on an orthorhombic unit cell ranging from 55.2 nm \times 58.9 nm \times 128.5 nm to 72.8 nm \times 76.3 nm \times 141.9 nm, and contained a solvent buffer of 10 nm in the (x, y, z) dimensions away from nonsolvent atoms. The AA potential energy function CHARMM36m (34, 35) for

proteins and the TIP3P (36) potential energy function for water were used. The AA systems were energy-minimized and equilibrated under constant pressure and temperature (NPT) conditions. Simulations in the constant NPT ensemble were performed using a Langevin thermostat at 310 K and a Nosé-Hoover-Langevin barostat at 1 atm. Bond lengths for hydrogen atoms were constrained using the SHAKE algorithm (37). An r-RESPA integrator was used with a timestep of 2 fs; long-range electrostatics were computed every 4 fs (38). Long-range electrostatics were calculated using the particle mesh Ewald algorithm, and short-range, nonbonded interactions were truncated at 12 Å (39). IP₆ molecules were parameterized using the CHARMM General Force Field (40). All simulations used the AAMD simulation package NAMD 2.14 (41). Production-level runs were performed on six capsid containing liquid water and two capsids containing IP₆, the RNP core, or both IP₆ and RNP. For all subsequent analysis AAMD trajectories were sampled at 0.04-ns intervals.

Strain Calculations. The centers of mass for every five amino acid residues in the CA domains were used to subsample the capsid structure. For each center of mass, a deformation tensor that describes the local deformation of a point particle in a system relative to its neighboring particles was calculated: $\mathbf{F}_{i,t} = \left(\sum_{j \in N_i^0} | r_{ji}^0 \rangle \langle r_{ji}^0 | \right)^{-1} \sum_{j \in N_i^0} | r_{ji}^0 \rangle \langle r_{ji}^t |, \text{ where } | r_{ji}^0 \rangle = \vec{x}_j^0 - \vec{x}_i^0 \text{ is the difference}$ between the Cartesian coordinates of neighboring particles *j* and *i* with respect to the reference configuration. Similarly, $|r_{ji}^t
angle$ is the difference in coordinates at time t, and N_i^0 is the set of particles in the local neighborhood of *i* in the reference configuration, within a cutoff distance of 10 nm. Note that the deformation tensor, $F_{i,t}$, can be determined by minimizing $\sum_{j \in N_i^0} |\langle r_{ji}^0 | F_{i,t} - \langle r_{ji}^t ||^2$. The reference configuration was the fitted atomic model of the fullerene capsid after CG relaxation. The per-particle Green-Lagrange strain tensor is then $\varepsilon_i = (F_i F_i^T - I)/2$, with volumetric and shear (von Mises) strain invariants given by $s_V = Tr(\epsilon)/3$ and $s_s = [Tr(\epsilon^2) - t]$ $Tr(\varepsilon)^2/3|^{1/2}$, respectively. Strain tensors were computed for each frame of the trajectories, until the invariants converged to an equilibrium value (e.g., $\sum_{i} s_{V,i}^{t+1} - s_{V,i}^{t} < 0.01$) at frame $t \sim 32$ ns. Strain patterns persisted for the remainder of the simulation time (see also *SI Appendix*, Table S1). Average strain magnitudes, $\langle | s_V | \rangle$, were measured using the strain corresponding to residue 110.

Core Volume Analysis. Core volumes of the capsid were computed using a Monte-Carlo sampling strategy. Cartesian coordinates for the C α atoms of the CA domains were monitored and used to subsample the capsid structure. For each iteration, points within the periodic simulation cell were randomly generated (\vec{x}_{MC}), and assigned to the capsid interior, if $\|\vec{x}_{MC}\| < \|\vec{x}_{C\alpha}\|$, where $\vec{x}_{C\alpha}$ is the position of the C α atom that maximizes the projection of the point onto the capsid structure ($\vec{x}_{MC} / \| \vec{x}_{MC} \| \cdot \vec{x}_{C\alpha} / \| \vec{x}_{C\alpha} \|$). A total of 10⁹ iterations were performed, and core volumes were calculated from the total fraction of points inside the capsid, accumulated for each 0.04-ns interval frame. Fluctuations from the mean volume (V - V) were measured in each capsid trajectory. Normalized fluctuation probabilities were obtained by binning the amplitudes with a 15-nm³ step size, and subsequently fit to Gaussian distributions. Statistical uncertainty was assessed by block averaging. For each distribution, the time series was divided into three blocks. The SD in fluctuation amplitudes for the three blocks is reported. Using three to five blocks gave qualitatively similar results. Fourier analysis was performed by computing the discrete Fourier transform on the mean free volume fluctuations (42).

CA Conformational Distributions. AAMD trajectories from production-level simulations of CA systems were subsampled at 0.04-ns intervals and aggregated. CA monomers in the capsid were aligned by minimizing the rmsd of the protein backbone atoms with respect to the crystallographic structure (PDB ID: 4XFX). Cartesian coordinates for the nonhydrogen atoms of CA were recorded, and the density of atomic positions $\rho(\vec{r})$ was calculated, using a hard-sphere van der Waals approximation on a discretized grid with a spacing of 0.5 Å imes 0.5 Å imes 0.5 Å as in (43). The 3D density map for the CA monomer was subsequently contoured at G = 0.5, 2.3, and 4.4 kcal/mol, where $G = -k_{\rm B} T \log \rho$. The parameter, ξ , was used to specify the distance between the centers of mass of the amino acid backbones of residue 21 and residue 22 on the pore helices (H1) of adjacent CA domains. The angle, ϕ , is defined as the angle between three centers of mass in the amino acid backbone, two of which were positioned on the CA NTD (residues 50 to 52, 78 to 80, and 128 to 130; residues 63 to 65, 144 to 146, and 58 to 60) and the third positioned at the adjacent CA CTD (residues 188 to 192, 166 to 170, and 196 to 200). (ξ, ϕ) values were recorded for each 0.04-ns frame and aggregated across each of the simulated capsid systems (CA, CA-RNP, CA-IP₆, and CA-RNA-IP₆). Successive contours for the (ξ, ϕ) distributions are given in units of $-\log p$, where p is the probability.

CG Simulations of Capsid Rupture. CG models of the capsid were generated from the AA models by computing the center of mass of each CA monomer at a resolution of 1 CG particle or "bead" per CA monomer. CA–CA interactions were added using a network of Morse bond effective potentials between the eight nearest-neighbor particles $[U_{Morse} = k (1 - e^{-\alpha(r-r_0)})^2]$, where the equilibrium bond distance, r_0 , was set to the distance found between the neighboring centers of mass, and the stiffness, α , was set to 2 Å⁻¹. Excluded volume interactions between CG particles used a soft-cosine potential (U_{Vol}), with $A = 100 \text{ kcal mol}^{-1}$ and cutoff $r_c = 75 \text{ Å}$. All CG MD simulations were performed in constant NVT conditions, with a periodic cell of (2,000 Å × 2,000 Å × 2,000 Å) and a Langevin thermostat at 300 K. CG particles with excluded volume interactions were added every 1,000 CG MD timesteps in a spherical region with radius 150 Å centered at the origin, until the capsid ruptured. All simulations were performed in LAMMPS and proceeded for 10⁶ CG MD timesteps.

Cryo-ET Imaging of Fractured Cores. Endogenous reverse transcription (ERT) starting with purified HIV-1 virions was performed as described (13). Samples of cores undergoing ERT for 4 h at 37 °C were mixed with equal volumes of 10-nm BSA Gold Tracer (Electron Microscopy Sciences), and 3.5-µL aliquots were applied onto glow-discharged Quantifoil grids (Electron Microscopy Sciences), blotted to near-dryness, and then plunge-frozen into liquid ethane. Cryotomograms were acquired using an FEI Titan Krios electron microscope operating at 300 kV and equipped with either a Falcon III camera or a K3/GIF with a slit width of 20 eV. Tilt series were collected using the data collection software Tomography 4.0 (FEI) with an angular range of -60° to $+60^{\circ}$, an angular increment of either 1 or 2°, defocus values of 5 to 10 µm, and a nominal magnification of x29,000 (Falcon III) or x33,000 (K3), which correspond to a pixel size of 2.92 (Falcon III) or 2.69 (K3) Å. Tilt series were aligned by using IMOD (44). Weighted back-projection or the simultaneous iterative reconstruction technique was used to reconstruct tomograms in IMOD.

Lattice mapping was performed as described (13) using as a search template a previously determined cryo-EM map of a HIV-1 CA hexamer (EMD-3465) downsampled to 25 Å. Each capsid surface was modeled and oversampled as a mesh, using surface modeling tools in the Dynamo software package. Between 2,000 and 3,000 subvolumes were exacted from the tomograms, each centered on points on the capsid surface with points distributed as evenly as possible and sampling the entire capsid surface as if it were intact. Multiple iterations of density matching and alignment of the search template with the subvolumes were performed in Dynamo, which converges to a lattice map containing the positions and orientations of hexamers on the capsid surface. Uncertainty in the lattice map was determined by the cross-correlation between each hexamer and the local density contained in the subvolumes (*SI Appendix*, Fig. S6). A total of 14 HIV-1 cores were imaged and mapped, and all tomograms containing cores were analyzed.

Local Order Parameter (χ) Analysis. Atomic models for CA hexamers were aligned with the Cartesian coordinates and Euler angles for each subtomogram in the cryo-ET density. For each structure, Ca positions were used to compute a per-particle order parameter, χ , which quantifies the separation in the capsid lattice. χ is defined as the number of neighboring particles (within a cutoff distance of 10 nm) of the same chemical identity in a CA monomer (e.g., residue number) $N_{\rm P}$, normalized by the average $N_{\rm P}$ for a given residue. χ values < 1 correspond to lattice separations that were higher than average for the capsid structure, whereas χ values > 1 correspond to lattice separations that were lower than average. The capsid with the highest degree of strain (CA-RNP) from the MD simulations was used to measure correlations between χ and $\textit{s}_{V}.$ The values of χ and \textit{s}_{V} were measured for the protein residue with the largest variation in lattice separation (residue 110). The total number of CA monomers that contained a particle with $\chi < 0.55$ or n_{CA} and the total number of particles with $0.75 < \chi < 0.9$ or n_{CG} were measured for each capsid structure.

Data Availability. All study data are included in the article and/or SI Appendix.

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