Electron cryo-tomography studies of maturing HIV-1 particles reveal the assembly pathway of the viral core

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ABSTRACT

To better characterize the assembly of the HIV-1 core, we have used electron cryo-tomography (ECT) to image infected cells and the viral particles cryo-preserved next to them. We observed progressive stages of virus assembly and egress including flower-like flat gag lattice assemblies, hemispherical budding profiles and virus buds linked to the plasma membrane via a thin membrane neck. The population of budded viral particles contains immature, maturation intermediate and mature core morphologies. Structural characteristics of the maturation intermediates suggest that the core assembly pathway involves the formation of a CA sheet that associates with the condensed ribonucleoprotein (RNP) complex. Our analysis also reveals a correlation between RNP localization within the viral particle and the formation of conical cores, suggesting that the RNP helps drive conical core assembly. Our findings support an assembly pathway for the HIV-1 core that begins with a small CA sheet that associates with the RNP to form the core base, followed by polymerization of the CA sheet along one side of the conical core towards the tip and then closure around the body of the cone.
During HIV-1 assembly and release the Gag polyprotein is organized into a signature hexagonal lattice, termed the immature lattice. To become infectious, the newly budded virus must disassemble the immature lattice by proteolyzing Gag, and then reassemble the key proteolytic product, the structural protein p24 (CA), into a distinct, mature hexagonal lattice during a process termed maturation. The mature HIV-1 virus contains a conical capsid that encloses the condensed viral genome at its wide base. Mutations or small molecules that interfere with viral maturation also disrupt viral infectivity. Little is known about the assembly pathway that results in the conical core and genome encapsidation. Here we have used electron cryo-tomography to structurally characterize HIV-1 particles that are actively maturing. Based on the morphologies of core assembly intermediates, we propose that CA forms a sheet-like structure that associates with the condensed viral genome to produce the mature infectious conical core.
INTRODUCTION

Late events in HIV-1 replication are profoundly dependent upon the biochemical, structural and enzymatic properties of the virus’ two poly-proteins Gag and Gag-Pol (Reviewed in (1)). The Gag poly-protein (p55gag) is the key structural determinant of the nascent viral particle, and its protein components include (in order from the N- to C-terminus) p17 (matrix, MA), p24 (CA), spacer peptide 1 (SP1), p7 (nucleocapsid, NC), spacer peptide 2 (SP2), and p6 (Fig 2A). The Gag-Pol poly-protein (p160gag-pol) is a product of a frame-shift in the ribosomal reading frame that occurs at the junction of SP2 and p6. p160gag-pol is produced at a ratio of approximately 1:20 of p55gag, and includes the products of viral pol including the viral enzymes protease (PR), reverse transcriptase (RT), and integrase (IN) (1).

Both Gag poly-proteins are targeted to the plasma membrane for assembly by basic residues in MA that bind to the plasma membrane-specific lipid, phosphatidyl inositol (4,5) bisphosphate. Membrane binding triggers the exposure of a myristoyl group modification at the poly-protein N-terminus that functions to stabilize the protein at the plasma membrane. (reviewed in {Sundquist, 2012 #105}). Targeting of the viral genome to sites of virus assembly is mediated by interactions between the viral RNA Psi-sequence and the zinc-finger motifs within the NC domain of the Gag poly-proteins. Assembly of a new viral particle is initiated when membrane associated Gag multimerizes as a result of protein-protein interactions among CA domains, and RNA-protein interactions between the viral genome and NC. These interactions drive poly-protein self-assembly and their consequent organization as a rigid hexagonal lattice, termed the “immature” lattice. Progressive assembly of the immature lattice at the cell surface induces membrane curvature and initiates formation of the
viral bud. Subsequent recruitment of the cellular endosomal-sorting-complexes-required-for-transport (ESCRT) machinery through the Gag late-domain motif in p6 leads to membrane fission and release of the viral bud as an immature, non-infectious particle (Reviewed in (1)).

Within the newly budded particle, the immature Gag lattice is arranged radially; while the p55Gag N-terminal MA is associated with the inner leaflet of the viral membrane, the C-terminal NC bound to the viral RNA extends toward the center of the particle (2, 3). Although the immature lattice is important to particle formation and budding, it does not represent the infectious form of the virus. Formation of the infectious particle requires that the immature lattice be disassembled by PR-mediated proteolysis of Gag at specific cleavage sites (Fig 2A; (1)). Eventual release of CA from the Gag poly-protein by PR leads to the assembly of CA into a mature conical-shaped core within the viral membrane (1).

Both mature and immature particles are structurally defined by distinct hexagonal lattices that are dependent upon the self-assembly properties of CA. However, CA utilizes very different domain contacts to stabilize the two lattices (4, 5), and, as a result, they have profoundly different structural characteristics (2, 6) and distinct functions during viral replication (1). In the immature lattice, polyprotein hexamers are stabilized by CA-CTD/CA-CTD interactions and SP1, while inter-hexamer contacts are mediated by CA-NTD/CA-NTD interactions. In contrast, mature lattice hexamers are stabilized by both CA-NTD/CA-NTD and CA-NTD/CA-CTD interactions (7). Inter-hexamer contacts in the mature lattice are the result of the formation of a CTD dimer interface between adjacent hexamers, and a hydrophobic trimer interface that mediates hexamic or pentameric assemblies of CA hexamers after final cleavage at the CA-SP1 site (4). Because the CA contacts that stabilize the immature and mature lattices are so different, it is expected that the assembly program for
the mature core requires the full-disassembly of the immature lattice followed by *de novo* reassembly of a mature CA lattice as a conical-shaped core (8).

Although the proteolytic steps that lead to disassembly of the immature lattice are biochemically and temporally well-defined (9), the process by which CA assembles into the mature core is poorly understood. Previous ECT studies of mature virions led to two distinct models for core assembly. The first model proposes that core assembly is initiated at the wide base of the capsid and proceeds to the narrow tip. This “base-to-tip” assembly model was proposed by Benjamin, *et al* (6) based on their observations that the capsid bases were often consistent with respect to size, curvature and distance from the membrane, but the capsid tips were less well ordered and sometimes unclosed. Based on these observations, the authors argued that capsid growth initiated at the structurally regular base and proceeded to the tip, sometimes failing to completely close the structure.

In a separate study on the ultrastructure of the mature HIV-1 virions (10), the authors noted that regardless of the virion diameter, the conical capsid extended across the full diameter of the viral particle. The authors therefore proposed a “tip-to-base” capsid growth model that initiated at the capsid narrow tip and grew until it encountered the opposite membrane, where its growth would be redirected to close the structure at the base. In this model, the membrane is predicted to be an essential factor for the formation of the conical core. In vitro CA assembly experiments show, however, that purified CA, under conditions of high salt, can assemble cones independent of membranes (11).

Computational simulations suggested a third model in which capsid assembly proceeds as the non-equilibrium growth of an elastic sheet (12). This model suggests that CA forms a sheet that precedes formation of the mature fullerene cone-shaped core, and that sheet
growth drives curvature of the structure until its edges join in space. The core then incorporates additional CA units until the structure closes, albeit with some defects and gaps as documented by ECT (13). These simulations could recapitulate not only the conical core structures, but also the full-spectrum of CA assemblies observed for diverse retroviruses.

Intermediate structures comprising the HIV-1 maturation pathway have been difficult to document due to the rapidity of the process. Previous studies have used mutagenesis to arrest maturation at specific PR cleavage steps (14, 15), while others have used small molecules to inhibit just the final PR cleavage step (8, 16). These studies provided structural information regarding the disassembly of the immature Gag lattice, but not the assembly of the viral core. Additionally, accumulation of intermediates may not fully reflect the maturation process due to potential dominant negative effects the intermediates can have on assembly (17).

New work described here avoids the limitations of previous studies by analyzing the maturation process while underway. To this end, we have imaged by ECT the released viral particles found assembling on the edge of and after release from infected cells, and have characterized the structures found within these particles. Because this sample is asynchronous with respect to the maturation process, we expect that all potential capsid structures representing the assembly pathway are present. As a result, we can characterize capsid assembly without the complication of off-pathway issues that might arise due to the use of drugs or the introduction of mutations to manipulate the maturation process. Morphological characterization and quantitation of the structures present in the maturing particles have allowed us to propose a model for HIV-1 capsid assembly from the initial release of the immature particle, to the formation of the mature, infectious, fullerene cone. Our model
proposes that viral core assembly proceeds asymmetrically and not as a uniform assembly from one end to another (i.e., base to tip, or tip to base). Instead, our results support an assembly pathway that begins with a small CA sheet that associates with the RNP at the base of the forming viral core. Growth then proceeds rapidly from the base along one side of the capsid structure until it reaches the membrane, which redirects capsid growth back toward the base and leads to the structure’s eventual closure.

METHODS

Virus construct. The proviral construct encoding pNLEGFP-BglI was generously provided by Irvin Chen (UCLA). To generate the virus, human embryonic kidney 293T cells (ATCC) were co-transfected with the proviral construct and the plasmid encoding the vesicular stomatitis virus glycoprotein (VSV-G) envelope (obtained from the AIDS Research and Reference Reagent Program, Division of AIDS, NIAID, NIH: pHEF-VSVG from Dr. Lung-Ji Chang). Culture supernatants were collected 36 h post transfection and subsequently filtered through a 0.45 micron filter. Particles were purified and concentrated by ultracentrifugation at 100,000 g through a 20% sucrose cushion. The virus pellet was resuspended in PBS, aliquoted and stored at -80°C until used for infection.

Infected cell preparation. Human umbilical vein endothelia cells (HUVECs, Lonza, Walkersville, MD) were seeded in 6-well plates containing carbon-coated gold EM finder grids (2 μm hole-size and 2 μm hole-spacing; Quantifoil, Jena, Germany) and cultured for 24 h in a humidified incubator maintaining 37 °C and 5% CO₂. Cells were infected with the VSV-G pseudotyped NLEGFPDelta BglIVprX. Infection was carried out by spinoculation at
800g for 1 h at 16 °C. Spinoculated samples were washed two times with fresh media prewarmed to 37 °C. Two ml of fresh media were then added to each well and samples were incubated for 36 h to allow for virus expression before cryo-preservation.

**Purification of “end-point” virus sample.** Viral particles were purified and concentrated from the culture supernatants of infected HUVECs. Briefly, HUVECs were infected as described above and culture supernatants were collected 36 hours post-infection and filtered through a 0.45 um filter. Filtrates were then placed back in the humidified incubator maintained at 37 °C and 5% CO2 for an additional 16h to allow purified particles to fully mature. Filtrates were then treated with 1 Unit TURBO DNase (Ambion) per 1 ml filtrate supplemented with 10 mM MgCl2 at 37C for 2 h. Viral particles were concentrated by ultracentrifugation at 100,000 x g over a 20% sucrose gradient, resuspended in HBSS and finally filtered through a 0.22 um filter.

**Electon cryo-tomography of HIV-1 infected whole cells.** EM grids on which virus-producing cells were growing were removed from the culture plate using forceps and treated with 3 μl of warm media containing 10 nm gold fiducials. Forceps and grid were transferred to the environment chamber of a Vitrobot Mark III (FEI) maintained at 37 °C and 80% R.H. Excess liquid was manually blotted from the grids on one side before plunging into liquid ethane. Cryo-preserved grids were then imaged in a 300-kV FEI G2 Polara, or FEI Titan Krios transmission electron microscope both equipped with a field emission gun and energy filter (slit width set at 20 eV). Data were collected with either a Gatan Ultracam 4kx4k lens-coupled charge-coupled device or a K2 Summit direct detector. Tilt series were collected.
over a series of angles ranging from -60 to +60 degrees using a step size of 1°; 22,500x magnification (effective pixel size of raw data is 5Å), a total dose of 150 e/Å², and a defocus of -6 μm. UCSF Tomo was used to collect the tilt series and 3-D reconstructions were carried out using a weighted back-projection algorithm tracking 10 nm fiducials in IMOD.

Gag assemblies, budding particles and budded virions (immature, maturing and mature) were identified based on ultrastructural features. Budded particles were classified as either immature, intermediate or mature based on the core morphology.

**Image analysis.** The Amira package was used to segment the electron density maps of the viral particles. Isosurfaces were generated using a thresholding approach to first select the most dense and prominent feature within the particles. Thresholding values were then adjusted to minimize noise while maintaining the feature and contiguous densities through multiple tomographic slices. From the resulting segmentations, an isosurface was generated for CA, the RNP and the viral membrane.

**Statistical analysis of RNP localization in viral particles.** In more than 60 tomograms, 1003 budded particles were identified and evaluated independently by three individuals. Each particle was assessed for the shape of its core and the location of the RNP within the viral particle, and immature particles were excluded from subsequent analysis. Core shape was characterized as one of three categories: cylindrical, conical, or pleomorphic, and the RNP location was identified using a set of X, Y, and Z coordinates to mark its centerpoint. The positional uncertainty of the RNP was calculated from the three coordinates provided by the three individuals, and only particles whose RNP uncertainty was within 28 nm (i.e., half the
distance of the reported core diameter of 56 nm (6) were considered further. We also limited our final analysis of RNP and capsid shape correlation to particles that had a single capsid structure that was either conical or cylinder shaped. Of the original particles, only 306 met all of these criteria. For the 306 chosen particles, the average X, Y, and Z coordinates that identified the position of the RNP in the tomogram were used to determine if the RNP was inside or outside of the core. The two-tailed p-value result reflects the probability of randomly observing a pattern of RNP location and core shape as observed in our data using 306 particles. The two-tailed p-value was 2.38 x 10^-6. Analyzing the data with Fisher’s exact test yielded a significant result (P<0.001). We conclude that there is a significant correlation between the shape of the core and the location of the RNP.

RESULTS

ECT of HIV-1 infected HUVECs

We used ECT to image viral particles that were frozen next to infected HUVECs growing on EM grids (Fig. 1A). HUVECs were selected for this study because we found that they are amenable to imaging by ECT; they attach well to the carbon surface that coats the EM grid, grow and divide on the EM grid in a manner indistinguishable from tissue culture plastic ware, and are readily infected by, and produce HIV-1 particles. Because the extreme edges of the HUVECs are very thin (<500 nm), these areas of the cells can be imaged by ECT, which is limited by sample thickness due to the high tilt angles that must be included for 3-dimensional (3D) volume reconstruction (18).
Multiple budding profiles demonstrating various stages of virus egress were observed at the plasma membrane (Fig. 1B). The immature gag lattice underlying the membrane in the budding profiles was readily identified based on the characteristic radial densities (Fig. 1B), as well as the hexagonal arrangement of the lattice in surface views (2, 3; Fig. 1E). In some cases, the same Gag lattice appeared on the surface of the cells as islands of flat sheets arranged loosely around a center point like the petals of a flower (Fig. 1D). To illustrate that the flat lattice was in fact assembled Gag, as opposed to clathrin which is also known to form flat lattices on the cell surface, we compared the Fourier transforms of the flat gag lattice (Fig 1E) with that of a clathrin coated vesicle (Fig. 1E compare left column with right column). Indeed, the lattice unit spacing of the two types of lattices were distinct (Fig. 1E). However, clear similarities were apparent when we compared the flat Gag lattice with the Gag lattice of the immature particle, again suggesting that the surface lattice is part of the HIV assembly program. While this “petal-like” pattern is somewhat unexpected, it may provide an explanation for the observed incompleteness of the Gag lattice in immature virions, and impart information regarding the earliest steps of virus assembly.

The budding profiles we observed at the thin edge of the infected HUVECs were primarily half-dome shaped (Fig.1C) suggesting that this configuration likely represents a kinetically slow assembly intermediate of the budding process. In contrast, more spherical budding profiles that appeared closer to egress and were linked to the cell surface by a narrow membrane neck (Fig. 1B) were less abundant. These observations are consistent with previously published reports (19, 20) indicating that Gag assembles on the surface of the cells over the course of 9-12 minutes and remains stably associated with the membrane before
components of the ESCRT machinery are recruited and more rapidly carry-out the membrane
scission event.

Collectively, these observations confirmed that the HUVECs were actively infected at the time of cryo-preservation, and that new viral particles were being continually released into the extracellular space.

Distribution of morphologies among particles in the non-purified virus sample

Because the particles observed here were not subjected to any purification process, but frozen and imaged directly in the culture environment in which they were produced (Fig 1A), we refer to this sample as an “in cellulo” sample. In total, we imaged 2121 in cellulo particles. For 91 of the particles we were unable to determine capsid morphology and therefore excluded these particles from subsequent analysis. As summarized in Table 1, the remaining 2030 particles were characterized as either mature (76%), immature (8%) or maturation intermediates (16%).

Mature viral particles in our sample demonstrated structural morphologies similar to those reported previously for purified particles (6, 10). Specifically, many of these mature particles contained the characteristic fullerene-cone-shaped capsid that extended across the full diameter of the particle membrane (Fig. 1A). Immature particles in the in cellulo sample were also morphologically similar to particles described previously (2, 3; Fig 1A). Of particular note was the observation that the immature lattice often did not fully cover the inner membrane surface of the viral particle (Fig. 1A, inset). These findings agree with previous reports (2, 3) suggesting that the immature Gag lattice is incomplete, but dispute the findings of Kol and colleagues (21). Collectively, these observations indicate that the mature and
immature particles from the *in cellulo* sample structurally match those found in purified preparations.

An important feature of the *in cellulo* sample that distinguished it from purified samples was the distribution of mature and immature morphologies within the particle population (Table 1). The *in cellulo* sample comprised approximately 76% mature particles, compared to approximately 94% reported previously for purified samples (14). Consistent with the observed decrease in the number of mature particles in the *in cellulo* sample was an increase in the relative number of immature particles. Previous reports indicated that just 3% of the viral particles in a purified population are immature (14), but our sample contained 8% immature particles. Furthermore, 16% of particles in the *in cellulo* population demonstrated morphologies that were neither mature nor immature, suggesting that these particles contain intermediate capsid structures formed during capsid assembly. The above observations suggest that as many a 24% (i.e., immature and intermediate particles) of the total *in cellulo* population were in the process of maturing at the time the sample was cryo-preserved.

In further support of our conclusion that the *in cellulo* sample was enriched for particles that were immature or maturing, we purified viral particles from the infected HUVEC cultures and allowed them to fully mature before freezing and imaging them by ECT. This purified sample showed a similar distribution of morphologies as that previously reported for purified particles (Table 1) (14), with the overwhelming majority (99%) of the particles being mature.

**Structural characterization of HIV-1 maturation intermediates**
We carefully analyzed the internal structures of the 333 particles in our *in cellulo* sample that were morphologically neither mature, nor immature, and therefore likely maturation intermediates, and noticed that they could be categorized into 2 groups of intermediate forms: “proteolytic” intermediates and “assembly” intermediates; with each group containing its own distinct sub-groups.

**Proteolytic intermediates**

**MA-CA-SP1 proteolytic intermediate:** Among the intermediate structures observed in the *in cellulo* sample were particles that had a single, unclosed electron dense layer that followed closely (within 10 nm) the inner curvature of the membrane (Fig. 2C). These particles constituted just 6% (20/333) of intermediate particles suggesting that they are very short-lived intermediates. A similar morphology was previously described for viral particles bearing mutations in Gag that arrested p55 processing after cleavage between SP1 and NC (Fig 2A; (14, 15, 22). Presumably, the structural features of this proteolysis mutant represent the particle following the early SP1-NC cleavage event that separates NC and RNA from the immature lattice. The MA-CA-SP1 protein remains associated with the membrane as a single, thick layer of density, while NC and the RNA condense to form the ribonucleoprotein complex, or RNP.

**CA-SP1 proteolytic intermediate:** Approximately 14% (48/333) of the intermediate particles in the *in cellulo* sample had a single density layer that was measurably thinner than that of the MA-CA-SP1 particles (5 nm for CA-SP1 versus 9 nm for MA-CA-SP1; compare panels in Fig. 2C and 2D). This thinner density layer approximated the curvature of the membrane, but it often failed to follow the membrane consistently. These characteristics are
similar to the structural features described for the Gag CA5 mutant (15, 22), as well as for particles treated with the maturation inhibitor Bevirimat (8, 16). In both cases, the final Gag cleavage event between CA and SP1 is inhibited and the mature capsid is not formed. The continuous thin single layer density is presumably the CA-SP1 protein that is stabilized by a bundle of six SP1 helices that initially form contacts in the immature lattice (2, 3). Failure to fully proteolyze the CA-SP1 prevents maturation and leads to a loss of infectivity (17, 23).

**Fully disassembled proteolytic intermediate:** The final class of particles that we grouped with the proteolytic intermediates included those particles that appeared to have no organized internal structures (Fig. 2E; Empty). Overall, these particles were more electron-dense than vesicles that were sometimes observed in the sample and were comparable in size to the viral particles, but could not be conclusively identified as virions. The existence of such “empty” particles would suggest that the immature lattice is at some point fully disassembled during the maturation process. However, these particles were also rare, comprising just 4% (13/333) of the intermediate particles, suggesting that they are either a very short-lived maturation intermediate, or not part of the normal HIV-1 maturation pathway.

**Capsid assembly intermediates**

In addition to the “proteolytic” class of maturation intermediates, we also observed particle morphologies that we classified as “assembly” intermediates (Fig. 3).

**Small Sheets:** This subclass comprised approximately 22% (72/333) of the intermediate forms observed, and included particles that contained small sheets of density (Fig. 3A). The particles in this subclass were structurally diverse with some of the sheets...
demonstrating high curvature (Fig. 3A, red arrows). Regardless of their shape, however, such densities were typically found near the membrane of the particles, and were characteristically thin. We distinguished this class of assembly intermediate from the putative CA-SP1 proteolytic intermediates because they were typically smaller than the CA-SP1 structures that arose as a result of mutation (15) or treatment with a maturation inhibitor (16). Additionally, CA-SP1 densities typically follow the gradual curvature of the membrane while the “small sheets” assembly intermediates demonstrated either a straight edge or a highly curved morphology—both distinct from the morphologies described previously for proteolytic intermediates (15, 16).

In some instances, the small sheet structures were associated with an electron dense feature that we interpreted as the RNP (Fig. 3A, yellow arrows). Because RNPs are not clearly defined, the location of putative RNPs was estimated independently by three individuals and only further considered if all three individuals identified the same location (within 25 nm of each other). Of the 43 maturation intermediates exhibiting small sheets, 9 also exhibited putative RNPs (identified in the same position by all three individuals). The average distance from the putative RNP center to the small sheet was 33 nm (±17 nm). When a similar analysis was applied to RNPs and conical capsids in mature particles from the same tomograms, the distance (i.e., from center of RNP to capsid layer at base) was found to be 29 nm (±6 nm). A non-parametric two-tailed t test (p=0.7567) indicated that the difference in averaged distances was not significant. Together this suggests that the RNP sometimes condenses and associates with capsid sheets soon after assembly begins.

Hooks: This intermediate subclass included the largest number of intermediate particles (Fig. 3B), with approximately 34% (112/333) of the particles that we classified as
intermediate comprising this category. Because the “hook” subclass represented such a large proportion of our assembly intermediates, we infer that exit from this sub-class is a slow step in the assembly pathway. Hooks also had a small curved sheet, but the signature structural feature of this sub-class was the presence of a single straight edge density that extended across the diameter of the particle from the curved end. The curved ends of these structures (Fig. 3B, red arrows) resembled the wide base of a conical mature core and often were associated with the RNP (Fig. 3B, yellow arrows). The similarity of these “hook”-like structures to the mature conical HIV-1 capsid was striking. Indeed, they appeared to be cones except that they were missing one wall and the narrow tip.

**Large Gaps:** This intermediate subclass accounted for 20% (68/333) of the intermediate particles and was defined by the presence of a clear cone structure with a major gap (Fig. 3C). The gaps observed for these particles were typically localized to one side of the cone, or near the cone tip consistent with earlier reports (6, 13). We consider the “large gaps” subclass to be more progressed assembly intermediates compared to the “hooks” subclass and therefore conclude that the final steps of the core assembly pathway involve closing the structure in the areas of high curvature.

**Intermediate structures are absent from the fully mature, purified virus sample.**

To confirm that the intermediate capsid assemblies observed in non-purified virus were “on-pathway”, and not simply aberrant capsids, we purified virus from infected HUVECs and allowed the purified particles time to fully mature before cryo-preservation and ECT. The purified virus sample contained primarily mature capsids that were either cones (71%, Table 1; Fig. 4A), or cylinders (15%, Table 1; Fig. 4B). Approximately 13% of the
purified particles showed an aberrant morphology (Table 1). Analysis of the aberrant particles suggested that they are irregular, closed structures (Fig. 4C). Importantly, the purified particles do not appear to contain capsid structures that are similar to the intermediate capsid morphologies (i.e., hooks and large gaps) described above.

Intermediate particles are less spherical than mature, purified particles.

Another characteristic that distinguished the in cellulo intermediate particles from the purified particles was that the in cellulo particles appeared less spherical, and in many instances demonstrated flat membrane edges (compare viral particles in Fig. 3 with those in Fig. 4). To quantify sphericity of individual viral particles, we used a simple ratio of length:width. We selected the particles in Fig. 3B and 3C for analysis as well as other intermediate and conical capsids found in the same tomograms. For each particle, we measured the length (greatest diameter through center of the particle) and width (diameter through center of the particle orthogonal to the length) and considered the ratio of those values (L:W). For spherical particles the L:W ratio is expected to be 1 and for non-spherical particles the ratio will be greater than 1.

We carried out the sphericity analysis for intermediate (n=37) and mature conical (n=37) particles from the in cellulo sample, and mature conical particles from the purified sample (n= 49). The results of this analysis showed that the intermediate particles from the in cellulo sample were the least spherical (L:W= 1.229), followed by the mature conical in cellulo particles (L:W= 1.150). The purified conical capsids were the most spherical (L:W= 1.118). Direct comparison (unpaired, two-tailed t test) of the average sphericity of the in cellulo intermediates with the in cellulo conical particles gave a result of p= 0.0337,
suggesting that the difference in sphericity is somewhat significant. The difference in sphericity observed for intermediate and purified conical particles was more significant ($p=0.0003$), supporting the idea that intermediate particles are less spherical than their mature counterparts. We also observed a difference in sphericity between the in cellulo conical and purified conical particles ($p=0.0183$). This result suggests that the process of purification (i.e., filtration, ultracentrifugation and resuspension) increases particle sphericity.

**RNP localization and mature capsid morphology.**

RNP formation precedes the completion of capsid assembly (15) and may help direct the assembly pathway (24). While the overwhelming majority (97%) of conical cores enclosed a density consistent with the features of the RNP (Fig. 5A), cylindrical cores enclosed RNPs just 67% of the time (Fig. 5B, Table 2).

**DISCUSSION**

The in cellulo particle population represents a unique experimental system to study the dynamic process of maturation. We used ECT to structurally characterize HIV particles that were maturing at the edge of infected human cells cultured on EM grids. The particles were cryo-preserved and imaged directly in the environment where they were produced and without additional purification. Because there is no purification step that would afford the particles time to fully mature before imaging, our sample contains particles at various stages of assembly, egress and maturation. Unlike previous studies that characterized viruses with mutations in cleavage sites of Gag (14, 15), or were treated with a maturation inhibitor (8, 16), our experimental design avoids the limitations of such strategies that often
produce aberrant capsids and non-infectious virus. As a result, we are able to describe maturation, and the authentic intermediates that define the process, as it occurs unaltered in newly budded viral particles.

While it is not possible to know if every intermediate structure observed is on-pathway to form a conical core, the overwhelming consistency in the “hook” and “large gap” structures, and their similarity to the fully mature conical capsid that defines the infectious HIV-1 particle, are compelling arguments that these structures are precursors to the mature capsid. Also, the relatively large number (75%) of non-mature/non-immature particles that display these morphologies (i.e., small sheet, hook, large gap) strongly suggest that HIV capsids often sample the assembly pathway(s) that lead to these structures, and then, often, to a mature conical core. We also note that the final distribution of mature capsid morphologies in our fully-mature, purified virus sample is comparable to what has been described previously ((10, 14); Table 1). This suggests that the HIV-1 maturation pathway within the \textit{in cellulo} sample yields HIV-1 capsids that are morphologically similar to previously described purified particles.

Furthermore, structural analysis of the mature endpoint viruses failed to identify similar intermediate structures among particles that were allowed to fully mature, and to our knowledge the “hook” and “large gap” structures have not been previously reported in the literature. We interpret these points to mean that the intermediate structures observed among the \textit{in cellulo} particles are not aberrant capsid forms that arise from a dead-end process, but instead they are likely structural intermediates that define the HIV-1 capsid assembly pathway. As a result of our unique experimental design, we are able to provide the first description of authentic HIV capsid assembly intermediates.
Maturation intermediate particles suggest that the HIV-1 capsid initially assembles as a sheet. The end-to-end models (i.e., tip-to-base, or base-to-tip) predict that capsid assembly intermediate forms will be dominated by structures that have either a tip but no base, or a fully formed base but not tip. In our sample, such open structures were not routinely observed. Importantly, the asymmetric shape of the “hook” intermediate structures argues against the end-to-end maturation models that describe core assembly as initiating from one of the extreme ends of the viral cone and proceeding circumferentially to the opposite end.

Instead, the morphologies of the assembly intermediates we observed support a capsid assembly pathway that is best described as a hybrid of the three previous models. We propose a model of assembly that begins with the polymerization of a small sheet of CA that associates with the RNP to form the base of the capsid structure. Continued growth of this small sheet from the capsid base progresses along one side of the cone in a manner similar to the simulations of Levandovsky and Zandi (12), which described capsid assembly as the non-equilibrium growth of an elastic sheet. Their simulations demonstrated the rapid growth of the conical shell along one side of its long axis in a manner that could lead to an intermediate structure that is very similar to the “hook” intermediate we observed in cellulo. Our images suggest that the sheet of CA then grows around the body of the cone to different degrees of completion (13).

The number of particles in each assembly intermediate subclass suggests relative kinetics of capsid assembly. The relative number of particles belonging to each of the assembly subclasses likely reflects the kinetics of assembly for the different intermediate structures. Our observation that the “small-sheet” subclass contains fewer particles than the
“hooks” subclass suggests that the initial growth of the capsid along one side of the cone down its long axis proceeds rapidly. The “hook” subclass is the most abundant assembly intermediate. Because these structures are typically a straight line density extending from the curved base, we suspect that the slower kinetics may be due to the need for the forming capsid to incorporate pentameric assemblies of CA at the highly curved base or tip in order to continue growing.

The viral membrane and RNP likely guide the core assembly pathway. In vitro, CA has the strong preference to assemble into tubes, spheres, and sheets, but rarely into cones (11). High salt conditions in vitro help promote the formation of cones (11), but such ionic strength does not reflect conditions found inside the virions in vivo.

In vivo factors that may promote formation of conical cores include the viral membrane and the RNP. The viral particles in our in cellulo sample were often less spherical than those particles in purified samples (compare Fig. 3 and Fig. 4). We propose that the apparent membrane deformations are not abnormal, but rather a function of the maturation process. Maturation may distort the viral membrane by forcing it to accommodate the growing, rigid capsid structure; for example, growth of a CA sheet along one side of the particle may cause that area of the particle membrane to become flat. Contact of the capsid with the viral membrane may not only affect membrane curvature, but also provide feedback to the growing mature lattice (6, 10). In those areas where the viral membrane is highly distorted by the capsid, the membrane may prevent continued assembly unless the capsid can adopt a localized curvature such that its growth is no longer impeded. Such curvature could be achieved, presumably, through the incorporation of pentamers at the narrow tip or wide base as predicted by the fullerene cone model.
Our observation that the assembly intermediate structures were often associated with a condensed RNP indicates that RNP formation precedes the completion of capsid assembly. Although we are not able to say conclusively when the RNP forms, early formation would suggest that the complex is important to capsid assembly and may bias the assembly pathway towards cones. Additionally, the strong correlation we observed between conical core formation and localization of the RNP to the interior of the capsid also suggests that the RNP influences the capsid assembly pathway. Indeed, such a role for the RNP in determining the structure of the HIV-1 capsid has been proposed (6), and it may be a mechanism of directed capsid assembly across diverse viruses (24). Collectively, these findings suggest that features of the RNP might represent potential therapeutic targets.

Other factors may also nucleate capsid assembly (11). In this case, capsid assembly would initiate independent of the RNP, but an assembly pathway that favors formation of a cone would preferentially enclose the RNP in the growing capsid structure. In contrast, a growing capsid that samples an assembly pathway that leads to formation of a cylinder would tend to exclude the RNP from the capsid interior due to the limited space within a uniformly narrow cylinder.

Model for HIV-1 capsid assembly. Based on our data, the HIV capsid assembly pathway (Fig. 6) is best described as an asymmetric assembly program mediated by the de novo formation of a CA sheet from dissociated CA subunits. We propose that a small sheet of CA initially forms and associates with the RNP. This CA sheet then grows rapidly along one side toward the narrow tip, followed by slower growth of the structure around the body of the cone. The inherent curvature of the CA sheet together with the RNP ultimately drives cone formation and inclusion of the condensed viral genome within the capsid interior.
ACKNOWLEDGEMENTS

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REFERENCES


Figure 1. Ultrastructure of HIV-1 infection in human umbilical vein endothelial cells (HUVECs). (A) Infected cell surrounded by newly released viral particles at various stages of maturation. Insets are enlarged views of an immature viral particle (green boxes), and a mature particle with a conical capsid (red boxes). (B) Late-stage HIV-1 budding profile on the surface of an infected cell. (C) Characteristic “half-dome” shaped early assemblies of immature Gag on the cell surface. (D) Flat Gag assemblies on cell surface. (E) Comparison of lattice features from the flat Gag assemblies (left, area from blue box region in D), an immature particle (center) and a clathrin coated vesicle right. Top panels are tomographic slices through respective lattices, middle panels are areas of the lattices (yellow box) that were compared by Fourier transform, results shown in bottom panels. Scale bars: A-D, 100 nm A-D; E, 10 nm.

Figure 2. Proteolytic intermediates formed during HIV-1 maturation. A) Schematic representation of the proteolysis steps that constitute the HIV-1 maturation process. B) Immature viral particle from the in cellulo sample. (C-E) Tomographic slices through viral particles that contain structures that represent the indicated maturation intermediate; (C) myr-MA-CA-SP1 polyprotein, (D) the last proteolytic intermediate, CA-SP1 and (E) viral particles with fully proteolyzed polyprotein. Scale bar 100 nm.

Figure 3. Assembly intermediates formed during HIV-1 maturation. (A-C) Tomographic slices through viral particles representing the indicated maturation intermediate. (A) Small – sheets, (B) “hooks” and (C) “large gaps.” Red arrows identify CA sheets (A) or the ends of the CA sheet forming the hook structure (B); yellow arrows, RNP. Scale bar 100 nm.
Figure 4. Capsid assembly intermediates are not observed in the fully mature, purified virus sample. Cryo-tomographic slice through HIV-1 particles purified from cultures of infected HUVECs with (A) a conical capsid, (B) a cylindrical capsid, or (C) an irregular capsid. Scale bar 100 nm.

Figure 5. Correlation between cylinder formation and RNP exclusion. Cryo-tomographic slice through HIV-1 particles with either (A) a conical shaped capsid that encloses the viral RNP, or (B-D) aberrant cylinder shaped capsids that failed to enclose the RNP (RNP indicated by yellow arrows). Scale bar 100 nm.

Figure 6. HIV-1 capsid assembly pathway. Tomographic slices through viral particles that contain intermediate capsid structures at various stages of maturation are ordered to suggest progressive assembly of HIV-1 capsid intermediate structures. Top row: Single tomographic slices through the particle volume showing the most complete view of the capsid structure. Middle row: Computational segmentation of the particle’s capsid density overlaid on the tomographic slice from top row. Bottom row: Full 3D segmentation of the entire particle volume. purple, membrane; green, immature capsid; blue, capsid assembly intermediate; red, mature capsid; yellow, condensed RNA genome.
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\[ p\text{-value} = 2.38 \times 10^{-6} \]
Figure 2.

A

B Immature: myr- MA CA SP1

C Type 1: myr- MA CA SP1

D Type 2: CA SP1

E Type 3: “Empty”
Figure 3.

A Small sheets

B Hooks

C Large gaps
Figure 4.
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Figure 6.