An atomic model of HIV-1 capsid-SP1 reveals structures regulating assembly and maturation

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Immature HIV-1 assembles at, and buds from, the plasma membrane before proteolytic cleavage of the viral Gag polyprotein induces structural maturation. Maturation is blocked by protease and maturation inhibitors (MIs), abolishing infectivity. The CA (capsid) and SP1 (spacer peptide 1) region of Gag is the key regulator of assembly and maturation, and the target of MIs. Here we applied optimized cryo-electron tomography and subtomogram averaging to resolve this region within assembled immature HIV-1 particles at 3.9 Å resolution and built an atomic model. The structure reveals a network of intra- and intermolecular interactions mediating immature HIV-1 assembly. The proteolytic cleavage site between CA and SP1 is inaccessible to protease. We suggest that MIs prevent CA-SP1 cleavage by stabilizing the structure, and MI resistance develops by destabilizing CA-SP1.

The major structural protein of HIV-1, Gag, oligomerizes at the plasma membrane of infected cells, leading to membrane bending and release of immature virus particles. Ordered cleavage of the Gag polyprotein at five sites by the viral protease (PR) then causes a dramatic structural rearrangement of the virus to the mature, infectious virion (fig. S1, A to C). Gag-Gag interactions in the immature virus are mediated by the CA domains and the adjacent SP1 region. After maturation, CA forms a conical core encapsulating the viral RNA-nucleoprotein complex (1). The final proteolytic cleavage in Gag occurs between CA and SP1. Even small remnants of uncleaved CA-SP1 have a dominant-negative effect on infectivity (2, 3). Cleavage at this site is inhibited by MIs (4) (fig. S1B), and polymorphisms in this region caused resistance against the first-in-class MI Bevirimat (BVM) (5). Other MIs inhibiting CA-SP1 cleavage are currently undergoing clinical trials, but the precise mechanism of their inhibitory action and of resistance is unclear.

Unprocessed Gag is assembled into irregular curved lattices whose structures cannot be determined using conventional structural biology techniques. The best structural model for CA within the immature virus has been derived by fitting available NMR and crystal structures of the folded N-terminal and C-terminal domains (CA-NTD and CA-CTD, respectively) into a structure of the immature CA lattice derived by cryo-electron tomography and subtomogram averaging (6). Subtomogram averaging can resolve protein structures within complex environments ranging from cells to enveloped viruses (7), but has been limited to resolutions of approximately 8 Å. The positions of α -helices, but not of amino acids, are visible at this resolution. Furthermore, 8 Å resolution is not sufficient to generate ab initio structural models for unknown structures such as the Gag regions downstream of the crystallized CA-CTD, which ends at residue 220 of the 231 amino acid CA domain (fig. S1D). This downstream region consists of amino acids thought to assemble a flexible hinge, followed by a sequence predicted to form a six-helix bundle that spans the C-terminal residues of CA, the cleavage site and the N-terminal residues of SP1 (8). This region contains the critical CA-SP1 cleavage site and includes the majority of amino acids essential for virus assembly (9-11). Obtaining a high-resolution structure of CA-SP1 in the immature arrangement is vital for a mechanistic understanding of HIV-1 assembly, maturation, and its inhibition.

The HIV-1 Gag construct \triangle MACANCSP2 (*11*) assembles into immature virus-like particles (VLPs) in vitro (Fig. 1A and fig. S1C). We assembled \triangle MACANCSP2 particles in the absence or presence of 100 µg/ml BVM. We also purified intact, immature HIV-1 virus particles carrying an inactivating PR mutation, D25A. For all three samples we acquired cryo-electron tomography tilt series implementing an optimized data collection scheme (*12, 13*) (table S1). VLPs with or without BVM were indistinguishable, showing densities for the Gag lattice similar to the immature virus particles (Fig. 1A). Subtomogram averaging was performed independently for each dataset using an optimized workflow implementing features including frame-based motion correction (*14*) and exposure filtering (*15*). The resolutions of the CA-SP1 layer in the final averages of untreated or BVM-treated VLPs, and immature virus were 4.5, 3.9 and 4.2 Å, respectively (Fig. 1B and fig. S2, A and B).

We compared the three structures and found no clear differences in the protein densities of CA or SP1 – they vary only in the presence or absence of densities at the center of the hexamer near SP1 (fig. S2, C to I, and fig. S3C). Given the high degree of structural similarity between the maps we used the 3.9 Å structure to build and refine a complete atomic model for the CA-SP1 region from Gag residues 148 to 371 (Figs. 1C and 2, figs. S3 and S4, table S2, and movies S1 and S2). Residues 356-371, covering the C terminus of CA, and the first eight residues of SP1, assemble into a six-helix bundle.

Interactions within the CA-NTD layer are described in fig. S4A. The CA-NTD and CA-CTD contact one another in the region of E160-E161 in the CA-NTD and Q308 in the CA-CTD. The relative positions of the two domains may also be restricted by the extended, rigid linker between them, which appears to be stabilized by an interaction with R305 in helix 8. Y277, the last residue of helix 7 in a solution structure (*16*) and in the mature-like CA hexamer (*17*), is rotated out of the helix, and packs against the linker at P279 (Fig. 2B). This network of interactions may allow the CA-CTD to be structurally modulated by cleavage upstream of CA-NTD and vice versa.

Highly conserved residues of the Major Homology Region (MHR) in CA-CTD abut the extended chain between the 3_{10} -helix and helix 8 (Fig. 2C). Charged residues in the 3_{10} -helix and extended chain make interactions with the neighboring CA-CTD molecule within the hexamer (Fig. 2, B and C). Hexamers are linked by a CA-CTD dimer interface (fig. S4B) formed by residues W316 and M317 (*6*, 18, 19).

Downstream of helix 11, beyond the residues that are resolved in crystal structures (*19*, *20*), the "hinge region" VGG (residues 353-355) unexpectedly adopts a rigid structure within the lattice. P356 then marks the start of a helix (referred to here as the CA-SP1 helix) that extends down to residue 371 in SP1 (Fig. 2, A and D) before abruptly ending, indicating that residues C-terminal of 371 are disordered (movie S1). The CA-SP1 helix protrudes up into the CA-CTD layer, where the top of the helix and the VGG "hinge" from one Gag molecule pack tightly against the CA-CTD from the neighboring Gag molecule. The CA-CTD, VGG hinge and CA-SP1 helix thus form a single, integrated "assembly unit" that defines the structure of the hexamer (Fig. 2, D and G).

This assembly unit appears to be stabilized by a three-

way interaction between H358 in the CA-SP1 helix, D329 in the base of helix 10, and P356 from the CA-SP1 helix in the neighboring CA molecule (Fig. 2E). K290 in the loop upstream of helix 8, and K359 in the CA-SP1 helix protrude from above and below these residues toward the center of the six-helix bundle where they coordinate a density, presumably a negatively charged ion cluster (Fig. 2F and movie S2). This arrangement is reminiscent of six arginine residues that protrude into the center of the NTD hexamer in mature HIV capsid (17, 21). 12 essential amino residues in CA-CTD have been identified where mutation to alanine abolishes virus assembly (22, 23). These are: W316 and M317 in the hydrophobic dimeric interface; V353, G354 and G355 in the VGG hinge; K290, D329, P356, H358 and K359, which together form the intricate network of interactions that define the assembly unit; and A360 and L363 which appear to make hydrophobic interactions within the six-helix bundle. There is therefore a remarkably close correlation between the sensitivity of a residue to mutation and a role for the residue in mediating interactions within the CA-SP1 assembly unit, confirming the importance of these interactions in virus assembly.

During maturation, the final proteolytic cleavage occurs between L363 and A364. In our structure, this site is in the middle of the CA-SP1 helix bundle where it is inaccessible to PR, which acts on extended protein chains (24) (Fig. 2D and movie S2). Disassembly of the immature lattice and full cleavage between CA and SP1 only take place once cleavage has occurred both between MA and CA, and between SP1 and NC (10, 25). Together, these observations support a model where the final step in maturation is regulated by limiting the access of PR to its substrate: cleavage upstream of CA and downstream of SP1 together destabilize the immature lattice and the CA-SP1 helix, thereby allowing PR to cleave between CA and SP1.

MIs block HIV infection by preventing cleavage at the CA-SP1 site (4, 13, 26) (fig. S5), and may also stabilize the immature lattice (27, 28). We mapped mutations, deletions and polymorphisms that confer resistance to BVM (5, 29, 30) and to another MI, PF-46396 (31, 32) onto our structural model (Fig. 3). This revealed that resistance mutations do not map out potential drug-binding pockets. Instead, they are located at protein-protein interfaces within the CA-SP1 lattice. Together with our observation that the CA-SP1 cleavage site is inaccessible in the immature virus, this implies that the mode of action of MIs is not steric inhibition of proteolysis but instead stabilization of the immature Gag lattice. Our data suggest BVM stabilizes the lattice by binding to a site in the center of the six-helix bundle (fig. S2). The reduced cleavage at the CA-SP1 boundary is a downstream effect of stabilization because cleavage of this site requires unfolding to an extended chain. HIV-1 appears to of MIs. aturation. Cold 12). Medline Medline

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develop MI resistance by destabilizing its immature form, thus directly counteracting the stabilizing effects of MIs.

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SUPPLEMENTARY MATERIALS

www.sciencemag.org/cgi/content/full/science.aaf9620/DC1 Materials and Methods Figs. S1 to S5 Tables S1 to S2 References (33–58) Movies S1 to S2

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Fig. 1. Structure of the immature HIV-1 CA-SP1 lattice at 3.9 Å.

(A) Computational slices through tomograms containing either immature HIV-1 or untreated or BVM treated Δ MACANCSP2 VLPs. Arrowhead marks the membrane bilayer in the left hand panel. Scale bar 50 nm. (B) Electron densities of CA-SP1 viewed perpendicular to the lattice, generated by subtomogram averaging from the samples shown in (A). One CA-SP1 monomer is highlighted in color with the CA-NTD and CA-CTD/SP1 in cyan and orange, respectively. The respective resolutions of the determined structures are annotated. (C) The refined atomic model shown in the same view as in (B) and rotated by 90°, viewed from outside of the virus. The 6-fold symmetry axis is annotated with a hexagon.



Fig. 2. Structural features in the immature CA-SP1 lattice. (A) A single CA-SP1 monomer as in Fig. 1C. α helices and the Cyclophilin A binding loop are annotated (α -helices from neighboring CA monomers are annotated in brackets). Colored rectangles indicate regions enlarged in (B) to (D). (B) The CA-NTD/CA-CTD linker is in an extended conformation with Y277 binding to the linker and S278 approaching R305. (C) The highly conserved residues in the MHR (Q287, E291, R299) stabilize the linker connecting the 310-helix and helix 8. Residues in this linker can interact with an adjacent CA monomer around the hexameric ring, e.g., R286 with E344, and D284 with R294; point mutations of these residues do not abolish assembly (22), suggesting some redundancy in these interactions. (D) The CA-CTD, VGG hinge, and the top of the CA-SP1 helix form an integrated structural assembly unit. The CA-SP1 cleavage site is marked by a blue asterisk. Dashed rectangles indicate approximate position of enlarged panels (E) and (F). Residues colored as in (E) and (F). (E) Residues D329, P356 and H358 (in purple) form a three way linkage between two neighboring CA-SP1 helices and the base of the CA-CTD. (F) K290 and K359 (in green) protrude from above and below the region shown in (E), to the center of the hexamer where they coordinate a strong density. (G) Horizontal (left) and vertical (right) slabs through the structure illustrate that the MHR (yellow), other residues in the CA-CTD base (red), the VGG hinge (blue) and the top of the CA-SP1 helix (pink) come together to form the hexameric assembly unit. The vertical slab shows one half of the hexamer represented in a surface view.



Fig. 3. Maturation inhibitor resistance mutations destabilize the immature lattice. Resistance mutations or naturally occurring polymorphisms that render HIV-1 resistant to BVM (red) or PF-46396 (green) have been mapped onto the atomic model. Mutations that confer resistance for both compounds are colored yellow. The CA-SP1 cleavage site is marked by a blue asterisk. For clarity only three helices of the six-helix bundle are displayed.



Editor's Summary

An atomic model of HIV-1 capsid-SP1 reveals structures regulating assembly and maturation Florian K. M. Schur, Martin Obr, Wim J. H. Hagen, William Wan, Arjen J. Jakobi, Joanna M. Kirkpatrick, Carsten Sachse, Hans-Georg Kräusslich and John A. G. Briggs (July 14, 2016) published online July 14, 2016

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