Preservation of HIV-1 Gag Helical Bundle Symmetry by Bevirimat Is Central to Maturation Inhibition

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ABSTRACT: The assembly and maturation of human immunodeficiency virus type 1 (HIV-1) require proteolytic cleavage of the Gag polyprotein. The rate-limiting step resides at the junction between the capsid protein CA and spacer peptide 1, which assembles as a six-helix bundle (6HB). Bevirimat (BVM), the first-in-class maturation inhibitor drug, targets the 6HB and impedes proteolytic cleavage, yet the molecular mechanisms of its activity, and relatedly, the escape mechanisms of mutant viruses, remain unclear. Here, we employed extensive molecular dynamics (MD) simulations and free energy calculations to quantitatively investigate molecular structure–activity relationships, comparing wild-type and mutant viruses in the presence and absence of BVM and inositol hexakisphosphate (IP6), an assembly cofactor. Our analysis shows that the efficacy of BVM is directly correlated with preservation of 6-fold symmetry in the 6HB, which exists as an ensemble of structural states. We identified two primary escape mechanisms, and both lead to loss of symmetry, thereby facilitating helix uncoiling to aid access of protease. Our findings also highlight specific interactions that can be targeted for improved inhibitor activity and support the use of MD simulations for future inhibitor design.

INTRODUCTION

The late stages of human immunodeficiency virus type-1 (HIV-1) replication involve the formation of particles that subsequently undergo maturation to generate infectious virions.1−4 Prior to maturation, group specific antigen (Gag) polyproteins are attached to the inner leaflet of the viral envelope in a pleomorphic, spherical, and quasi-hexagonal lattice.4−5 HIV-1 protease cleaves Gag between its domains during maturation, in which the rate-limiting cleavage resides between the capsid (CA) and spacer peptide 1 (SP1).6 In the mature infectious particle, the released CA molecules assemble into a cone that encloses the viral nucleoprotein complex. The body of the cone is formed by CA hexamers that pack with quasi-hexagonal symmetry, and canonically, the wide and narrow ends of the cone are capped with seven and five CA pentamers, respectively.7

Inhibition of HIV-1 maturation is the target of a novel class of small molecule drugs. Bevirimat (BVM), or 3-O-(3′,3′-dimethylsuccinyl)betulinic acid, was the first proposed maturation inhibitor and acts by blocking cleavage at the helical junction between the C-terminal domain of CA (CA_{CTD}) and SP1.8 However, clinical trials were halted due to loss of susceptibility in over half of patients.12,13 In vitro studies have shown that point mutations throughout the CA/SP1 junction confer BVM resistance, suggesting that the polymorphism in this region, and in particular in the QVT motif of SP1, affects the mechanism of action of BVM.14−16

The current mechanistic model for BVM proposes that it stabilizes the CA/SP1 junction. Recent structural studies using X-ray crystallography,9 cryoelectron tomography (cryo-ET),10 and electron diffraction of frozen-hydrated 3D microcrystals (MicroED)18 resolved this junction as a six-helix bundle (6HB). Intriguingly, the six scissile bonds (between L363 and A364 in each protomer) face the interior of the bundle in which the central pore sterically occludes access of HIV-1 protease. Cryoelectron microscopy (cryo-EM) and magic angle spinning nuclear magnetic resonance analysis of CA/SP1 protein tubules, which adopt mature-like conformations, suggest that the CA/SP1 junction exists in a dynamic helix–coil equilibrium in the mature state.19 Cryo-ET analysis of a series of Gag cleavage mutants similarly showed mature lattices with disordered CA/SP1 junctions.20 However, immature lattices maintained ordered, helical CA/SP1 junctions.20 It is therefore unclear whether immature Gag lattices undergo a dynamic helix–coil equilibrium and if this equilibrium is necessary for protease activity and, relatedly, affected by BVM.

The recent high-resolution (2.9 Å) MicroED structure of BVM-bound CA_{CTD}/SP1 revealed that a single BVM molecule stabilizes the 6HB via both electrostatic interactions with the dimethylsuccinyl moiety and hydrophobic interactions with the pentacyclic triterpenoid ring of BVM.18 Interestingly, a comparison between the drug-free and BVM-bound structures
suggests that no major structural changes throughout the 6HB are induced by BVM. Likewise, recent solid-state nuclear magnetic resonance (ssNMR) experiments confirmed that BVM-induced structural changes are subtle, which further suggested that there are minimal changes in the conformation of the 6HBs. From these experiments, it can be inferred that the 6HBs may not exist in a dynamic helix−coil equilibrium. Consequently, the molecular mechanisms driving BVM activity and, relatedly, how escape mutants recover viral activity remain unclear. More recently, cryo-EM and MD analysis suggested that the edges of the fissures within immature HIV-1 capsid particles expose incomplete Gag hexamers, which could be the potential substrates for protease cleavage. This would imply that only the protomers within incomplete 6HBs may be in a dynamic helix−coil equilibrium.

Given our understanding of the chemical interactions between BVM and CACTD/SP1, we sought to further address the detailed mechanism of action of BVM by performing all-atom molecular dynamics (MD) simulations of BVM bound to wild type (WT) Gag CACTD/SP1 and three CACTD/SP1 mutants that impart varying degrees of drug resistance, L363F, A364V, and V370A. The aggregate MD statistics (across 120 trajectories) encompassed 72 μs. We note that both the equilibrated position of the BVM pentacyclic triterpenoid moiety (see Figure S1) and the average WT 6HB structure (see Methods) were initialized: the dimethylsuccinyl moiety pointing toward the K359 ring (or “BVM-up”) and pointing toward the C terminus (or “BVM-down”). We simulated three configurations of WT virus (drug-free, BVM-up, and BVM-down), both with and without IP6, each with five independent MD replicas of 0.60 μs. We used the same procedure to simulate the escape mutants L363F, A364V, and V370A. The aggregate MD statistics (across 120 trajectories) encompassed 72 μs. We note that both the equilibrated position of the BVM pentacyclic triterpenoid moiety (see Figure S1) and the average WT 6HB structure sampled by our MD simulations (see Figure S2) are in agreement with experimental predictions. To characterize the 6HB conformations sampled across the WT and mutant viruses, we used a combination of dimensional reduction techniques. We described each 6HB configuration by a high-dimensional set of quantitative descriptors, a procedure known as featurization (see Methods for details).

Characterization of the Six-Helix Bundle Conformational Ensemble. We first analyzed the impact of BVM binding on the conformational ensemble of the 6HB. We prepared MD simulations of a solvated CACTD/SP1 hexamer (PDB code 5L93) with restraints to helix 9 in order to emulate the quaternary structure of the immature lattice (see Methods). As seen in Figure 1, BVM was positioned in the center of the 6HB. Two binding poses suggested on the basis of our prior 2.9 Å MicroED structure were initialized: the dimethylsuccinyl moiety pointing toward the K359 ring (or “BVM-up”) and pointing toward the C terminus (or “BVM-down”). We simulated three configurations of WT virus (drug-free, BVM-up, and BVM-down), both with and without IP6, each with five independent MD replicas of 0.60 μs. We used the same procedure to simulate the escape mutants L363F, A364V, and V370A. The aggregate MD statistics (across 120 trajectories) encompassed 72 μs. We note that both the equilibrated position of the BVM pentacyclic triterpenoid moiety (see Figure S1) and the average WT 6HB structure sampled by our MD simulations (see Figure S2) are in agreement with experimental predictions. To characterize the 6HB conformations sampled across the WT and mutant viruses, we used a combination of dimensional reduction techniques. We described each 6HB configuration by a high-dimensional set of quantitative descriptors, a procedure known as featurization (see Methods for details). Using all 120 trajectories (i.e., across all replicas for WT and mutant virus), we computed gyration moments for each ring of residues along the 6HB, i.e., each ring consisted of the same six residues, for each sampled configuration. These features were subsequently used for dimensional reduction.

Dimensional reduction was performed using time-lagged independent component analysis (tICA), a linear projection technique that aims to maximize representation of characteristic of maturation inhibitor activity rather than binding affinity to the 6HB pore.

### RESULTS

#### Characterization of the Six-Helix Bundle Conformational Ensemble

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autocorrelations (see Methods). The largest eigenmodes correspond to the slowest varying changes within the constraint of linear transformation of the feature set and have been used to describe collective variables of biomolecules. Here, we find that the first two tICs capture 25% of the kinetic variance. It is therefore evident that the 6HB conformational space is not easily projected onto a simple 2D manifold in tIC space. Instead, we find that the first 10 components (tICs) capture up to 84% of the kinetic variance. We then clustered the 10D tIC-projected data using k-means clustering and filtered the data to 7 clusters with the highest populations, representing 76% of all configurations (see Methods). For visualization purposes, the aggregated data projected onto these 10 tICs were embedded onto a 2D manifold using multidimensional scaling. Figure 2A depicts the filtered and labeled tIC data on the embedded 2D manifold.

Each of the 7 filtered clusters represents a distinct subpopulation of 6HB states, which were analyzed further. For each configuration within each cluster, we computed the root-mean-squared deviation (RMSD) of the 6HB Cα backbones with respect to a prior atomic model (PDB code 5L93)37. We also quantified the symmetry throughout the 6HB using the 2D 6-fold Mermin order parameter ($\psi_{6}^{2D}$) for each Cα ring along the 6HB (see Methods). In other words, $\psi_{6}^{2D}$ measures the average hexatic order on a per-ring basis throughout the 6HB; $\psi_{6}^{2D}$ is expected to be a value of 1 if the 6HB is 6-fold symmetric and diminishes toward 0 with increasing asymmetry. Finally, we quantified the similarity of the Cα backbone to an ideal $\alpha$-helix ($\alpha_{\text{sim}}$). When the 6HB is $\alpha$-helical, $\alpha_{\text{sim}}$ is expected to be 1 and diminishes toward 0 with increasing dissimilarity.

Figure 2. Clustering of six-helix bundle (6HB) configurational states. (A) Two-dimensional projection of the aggregate 10-dimensional time-lagged independent components using multidimensional scaling (MDS). Data points are colored according to their cluster index from k-means clustering. A randomly selected subset ($n = 7200; 2\%$ of data set) of the data is shown here for clarity. (B) Comparison of the 2D 6-fold Mermin order parameter ($\psi_{6}^{2D}$) and root-mean-squared deviation (RMSD) of all 6HB configurations within each cluster; cluster labels were sorted in decreasing order of $\psi_{6}^{2D}$ ($n = 360,000$). (C) Representative snapshots of the 6HB structure classified by each cluster. Each cluster shows top and side views of 10 randomly selected and aligned configurations with each helix colored separately. The percentage of the total population of the data that is represented by each respective cluster is shown in parentheses.
Figure 2B compares the $\psi_6^2D$ and RMSD across the 7 clusters, which were sorted with respect to the mean $\psi_6^2D$. We find that decreasing $\psi_6^2D$ across clusters is commensurate with increasing RMSD; i.e., the C1, C2, and C3 clusters correspond to the most symmetric conformations. However, $\alpha_{\text{sim}}$ remains around 0.96 in all clusters (see Figure S3). These results suggest that the 6HB remains largely $\alpha$-helical and fluctuations from the average 6HB structure, i.e., cluster 1, are largely due to distortions to the 6HB symmetry.

Representative configurations from each cluster are depicted in Figure 2C. Cluster 1 contains the most similar structure to the atomic model of the 6HB, which is both $\alpha$-helical and 6-fold symmetric. Clusters 2—4 are partially 6-fold symmetric. Here, loss of 6-fold symmetry is due to a combination of helical splaying (e.g., cluster 2) and contraction along lower-
symmetry axes (e.g., cluster 4). Additional losses of 6-fold symmetry are seen in clusters 5−7, which are ascribed to the disassociation of knob-in-hole interactions between adjacent helices. In the sections below, we analyze the role of BVM, IP6, and escape mutants on the expression of these clusters.

**Bevirimat Preserves while Escape Mutants Disrupt Symmetry in the Six-Helix Bundle.** Using our cluster definitions, we now quantify cluster populations in WT HIV-1 viruses, in which we compare both BVM-up and BVM-down binding orientations (also see Table S1). As seen in Figure 3A, the cluster with the majority population (89%) in WT virus without BVM is cluster 3, i.e., 6HBs with partial 6-fold symmetry; the minority population (11%) is cluster 6, i.e., asymmetric 6HBs. Interestingly, cluster 1, i.e., 6HBs with 6-fold symmetry, is negligibly populated in drug-free WT virus. The presence of BVM in both orientations, however, broadly shifts the cluster populations in favor of clusters 1 and 2. In the BVM-up orientation, the population of cluster 1 is 44% and more populous than that of the BVM-down orientation (27%). The primary difference between the two binding poses is the association between the dimethylsuccinyl moiety and the K359 ring in the BVM-up case, which was previously proposed on the basis of MicroED structural studies. This stabilization occurs even though the distance between the carboxylate of dimethylsuccinyl and the ammonium of K359 is greater than is typically seen for salt bridges, which may arise due to the stoichiometry of six lysines to one dimethylsuccinyl in BVM. Our analysis suggests that this interaction is productive for 6-fold symmetry preservation in the 6HB, but both poses appear to be viable binding modes and each restricts the loss of symmetry in the 6HB.

The three mutant CA<sub>CTD</sub>/SP1 hexamers exhibited varying levels of distortion to the 6HB in the absence of BVM. As seen in Figure 3A, taking increasing cluster indices as indicative of increasing loss of symmetry, we find that the preservation of 6HB structural integrity in the majority population varies approximately as WT > V370A > L363F > A364V. Note that all three mutant viruses express replication-competent phenotypes, suggesting that the observed fluctuations in the 6HB structure do not impede viral assembly and maturation. However, the increased flexibility in the 6HB, most apparent in A364V, may facilitate protease access to the CA/SP1 scissile bond. Indeed, in the case of A364V, enhanced proteolytic cleavage of CA/SP1 compared to WT has been observed.

By contrast, L363F and V370A had no discernible effects on proteolytic cleavage efficacy. The addition of BVM has both mutant-specific and binding-pose specific effects on cluster populations. The BVM-up pose does not significantly increase the population of 6-fold (cluster 1) or partially 6-fold (clusters 2−4) symmetric clusters in the three mutant viruses. Instead, a broad distribution is observed with the majority population of L363F, A364V, and V370A 6HB states in low-symmetry clusters (clusters 5−7). These results suggest that the BVM-up pose is unable to recover the high-symmetry states that are stabilized by BVM in WT virus and instead may have an antagonistic effect. However, the BVM-down pose shifts the cluster populations of L363F and V370A toward clusters 2 and 4, i.e., partially 6-fold symmetric states. The implication would be that the BVM-down pose may partially recover the activity of BVM seen in WT virus. In the A364V case, however, the majority population remains in clusters 5−7, i.e., low-symmetry states. Both BVM binding poses appear to be ineffective against A364V.

Prior assays have shown that the fold change (FC) in half-maximal effective concentration (EC<sub>50</sub>), i.e., the ratio of EC<sub>50</sub> in mutant viruses to that of WT virus, is 20−120 for V370A and >1000 for A364V. In separate CA/SP1 processing assays in the presence of BVM, the majority of CA/SP1 molecules were cleaved in A364V mutants, intermediate levels were cleaved in L363F and, to a lesser extent, V370A mutants, and small fractions were cleaved in WT virus. These assays imply that the experimental efficacy of BVM in blocking protease activity varies as WT > V370A > L363F > A364V. On the other hand, our cluster analysis implies that BVM-induced symmetry preservation varies as WT > L363F > V370A > A364V. Therefore, the ability of BVM to restrict the occurrence of low-symmetry 6HB configurations correlates with the efficacy of the drug, with the exception of L363F compared to V370A. We hypothesize that additional factors may influence BVM efficacy, and we explore one such possibility next.

**Inositol Hexakisphosphate (IP6) Has Cooperative Effects with BVM on Six-Helix Bundle Stabilization.** IP6 is an essential cofactor in HIV-1 replication and has been shown to bind to the aforementioned K359 ring in immature WT virus. Prior MD simulations indicate that IP6 has a stabilizing effect on the 6HB, similar to the observed effects of BVM. It is therefore of interest to investigate potential cooperative effects when both IP6 and BVM are present.

According to our cluster analysis, as seen in Figure 3B, the addition of IP6 to WT virus in the absence of BVM increases the population in majority cluster 3 by 7%. However, cluster 1 remains unpopulated, indicating that IP6 itself does not induce the same degree of symmetry preservation as BVM by itself (as seen in Figure 3A). The addition of both IP6 and BVM, however, increases the population in cluster 1 to 74% and 47% for BVM-up and BVM-down, respectively, an increase of 30% and 20% compared to BVM by itself. The significant increase in the cluster 1 population suggests that the presence of IP6 and BVM within the central pore of the 6HB cooperatively promotes 6-fold symmetric states. It is also interesting that this effect is more pronounced in the BVM-up case. In this orientation, the dimethylsuccinyl moiety interacts with the K359 ring that also interacts with IP6. Rather than competing, both molecules appear to share the K359 binding sites, with the dimethylsuccinyl moiety of BVM and IP6 coordinating 1−2 and 4−5 lysine side chains, respectively (see Figure S4), with BVM also tending to rotate within the interior of the 6HB (see Figure S5).

Several mutant-specific and binding-pose specific effects on cluster populations also emerge when IP6 and BVM bind to the 6HB. In the absence of BVM, both L363F and A364V mutants populate lower symmetry states, primarily clusters 4 and 6, while the V370A mutant primarily populates cluster 3, the same as WT virus. Hence, IP6 has varied effects on mutant viruses (Figure 3B in comparison to Figure 3A): IP6 increases 6HB symmetry preservation for V370A while decreasing the same for L363F and A364V mutants. However, the resultant trend in 6HB symmetry preservation remains the same as IP6-free virus: WT > V370A > L363F > A364V.

All three mutants exhibit greater symmetry preservation in the 6HB when BVM is added to IP6-bound virus, although to varying degrees and more substantially in the BVM-up pose. The V370A mutant populates cluster 1 with 58% for the IP6 and BVM-up case, while the L363F and A364V mutants populate cluster 2 with around 58% and 27%, respectively. For...
comparison, the WT virus populates cluster 1 with 74% probability, indicating a loss of symmetry upon mutation, which is to be expected. Taken together, these results show that the efficacy of symmetry preservation by IP6 and BVM-up varies as WT > V370A > L363F > A364V, which is consistent with prior FC-EC50 measurements and CA/SP1 processing assays.14,15,30,32,35 Note that the former two mutants exhibit greater ordering compared to BVM-free WT virus while the latter exhibits greater disordering. These general trends are preserved in the BVM-down case, but the specific clusters that are populated by each mutant tend to be those with lower symmetry compared to the BVM-up case. Importantly, our analysis shows that IP6 cooperatively affects the activity of BVM, and its role in experimental assays should be clarified in future maturation inhibitor studies.

**Escape Mutants Exhibit Multiple Mechanisms of Failure.** Having analyzed the effect of escape mutants on cluster populations with varying degrees of 6-fold symmetry, we now investigate the origin of these observed differences. In particular, we quantify intermolecular contacts between BVM and hydrophobic residues along the interior cavity of the 6HB, as depicted in Figure 4A. We narrow our analysis to clusters 1, 2, 4, and 5 to provide a representative selection of 6HB configurational states upon BVM binding; additional data from the remaining clusters are shown in Figure S6. In the following, we analyze conditions in which each of these clusters is highly populated, i.e., in the presence of BVM in the BVM-up orientation; IP6 is also included for clusters 1, 2, and 4.

Using the cluster labels assigned to each configuration, we aggregated ensembles of each of the aforementioned clusters across WT and mutant virus trajectories for further analysis. Figure 4B,C shows the computed probability distributions \( P(r) \) of the radial distance \( r \) between the C atoms throughout the highlighted rings in the pentacyclic triterpenoid motif of BVM and Cα atoms of L/F363 and M367 within the listed cluster index (i.e., for clusters 1, 2, 4, and 5). The dashed line indicates the mean \( r \) between the respective moieties from the atomic model derived by MicroED.18 (D) Snapshots showing representative positions of BVM (stick representation) within the central pore of the 6HB (ribbon representation).

![Figure 4](https://doi.org/10.1021/jacs.1c08922)
To further quantify the increases uncoiling propensity, behavior is seen in the cluster 2 and 3 distributions. The broadening of the distribution indicates that the L/F363 coordination around M367 value. The r367 distribution exhibits multimodal with the largest peak shifted to smaller ABsim = 14 and ψ2D = 0.60, which transitions to an uncoiled state that broadly spans a range of 4 < ABsim < 12 with a free energy barrier height of 2.32 ± 0.21 kcal/mol (see Figure 5B). Note that the uncoiled state also spans a broad range of 0.35 < ψ62D < 0.70 with a local free energy minimum around ψ62D = 0.50. Hence, helical unfolding is commensurate with a reduction in 6HB symmetry. As seen in Figure 5C, the addition of BVM to WT virus with IP6 introduces another basin representing the folded state (ABsim = 14) in a high-symmetry 6HB (ψ62D = 0.9). For uncoiling, two structural transitions occur in sequence. The 6HB first distorts into a lower symmetry state (ψ62D = 0.40) with a barrier height of 3.51 ± 0.48 kcal/mol, followed by

BVM uniformly interacts with L363 throughout the six helices, BVM may preferentially interact with a few M367 residues throughout the six helices, which is likely due to favorable hydrophobic interactions.

For cluster 2 (green lines in Figure 4B,C), the r367 distribution is multimodal with the largest peak shifted to smaller r compared to the experimental r367 value. The r367 distribution, on the other hand, remains largely unimodal. In this cluster, BVM appears to closely interact with a subset of the helices around M367 while remaining uniformly coordinated by L/F363; in other words, BVM tilts such that the bottom of the pentacyclic moiety establishes closer contact with the 6HB (see Figure 4D). For cluster 4 (gray lines in Figure 4B,C), the r363 and r367 distributions exhibit modularity with peaks appearing below the experimental r363 and r367 values. Here, the entire pentacyclic moiety of BVM appears to establish close contacts with a subset of helices around L/F363 and M367. Recall that clusters 2 and 4 both represent 6HBs with partial 6-fold symmetry, suggesting that BVM fails to establish close contacts with a subset of helices around L/F363 and M367. Hence, helical unfolding is commensurate with a reduction in 6HB symmetry.

As seen in Figure 5C, the addition of BVM to WT virus with IP6 introduces another basin representing the folded state (ABsim = 14) in a high-symmetry 6HB (ψ62D = 0.9). For uncoiling, two structural transitions occur in sequence. The 6HB first distorts into a lower symmetry state (ψ62D = 0.40) with a barrier height of 3.51 ± 0.48 kcal/mol, followed by

Loss of 6-Fold Symmetry in the Six-Helix Bundle Increases Uncoiling Propensity. To further quantify the effect of BVM, we compute free energy surfaces (FESs) that describe the helix-to-coil transition in the 6HB. Here, we use well-tempered metadynamics, a technique that dynamically adds external bias to the system to promote rare event sampling, from which the FES is recovered through reweighting (see Methods). We project the FES along two coordinates, ψ62D and the so-called α-β similarity (ABsim), which assesses the collective deviation of ϕ and ψ torsional angles from that of α-helices. When the torsional angles throughout the measured protein backbone are equivalent to α-helices, ABsim is expected to be 14 (twice the number of residues) and linearly diminishes toward 0 with increasing deviation, i.e., uncoiling.

We compare the FES computed for WT virus with IP6 (Figure 5A,B) to that of WT virus with IP6 and BVM (Figure 5C,D) and L363F mutant virus with IP6 and BVM (Figure 5E,F). Here, we focus on the L363F mutation, as our previous analysis suggests that the A364V mutation is effective due to destabilization of the 6HB, which BVM is unable to mitigate. In WT virus with IP6, two primary free energy basins are observed. The first is the folded α-helical state around ABsim = 14 and ψ62D = 0.60, which transitions to an uncoiled state that broadly spans a range of 4 < ABsim < 12 with a free energy barrier height of 2.32 ± 0.21 kcal/mol (see Figure 5B). Note that the uncoiled state also spans a broad range of 0.35 < ψ62D < 0.70 with a local free energy minimum around ψ62D = 0.50. Hence, helical unfolding is commensurate with a reduction in 6HB symmetry.
uncoiling with a second barrier height of 3.32 ± 0.39 kcal/mol (see Figure 5D). The distortion of the 6HB represents a significant barrier against uncoiling introduced by BVM. In contrast, this high-symmetry state is abolished in the L363F mutant with IP6 and BVM, as seen in Figure 5E. In this case, note that the 6HB tends to adopt low-symmetry configurations similar to that of WT virus. The barrier height for uncoiling, however, is 3.05 ± 0.35 kcal/mol (see Figure 5F), an increase of 0.73 kcal/mol compared to that of WT virus (p-value of 0.006 according to the unpaired t test), suggesting that the presence of BVM slightly decreases the propensity for uncoiling despite the symmetry loss throughout the 6HB. Nonetheless, it is clear from our analysis that BVM primarily prevents helical uncoiling by stabilizing high-symmetry 6HB states.

**DISCUSSION**

Taken together, our analysis suggests that BVM coordinates the 6HB in WT virus, thereby abrogating helical uncoiling, inhibiting protease access, and preventing maturation. To effectively and uniformly coordinate the 6HB, BVM stochastically rotates within the bundle, hopping from protomer to protomer (see Figure S5). Our simulations also suggest that both BVM-up and BVM-down are viable binding poses, although we predict that BVM-up is the more effective inhibitor; in the future, it would be interesting to investigate the relative population of these two binding modes in virions using computational techniques, such as the coarse-grained modeling and simulation framework previously used to study HIV-1 assembly.38–41 In mutant viruses, however, we observe two primary structural states that are related to their escape mechanisms and provide the following interpretation that is depicted using helical wheel diagrams in Figure 6.

First, escape mutations can disrupt the hydrophobic binding interface along the interior of the 6HB (cyan box in Figure 6), i.e., L363 and M367. According to the cluster 1 distributions in Figure 4B,C, BVM interacts more strongly with L363 compared to M367. Given the larger hydrophobicity of phenylalanine compared to leucine,42,43 the L363F mutant likely increases the microscopic binding affinity of BVM. Yet, due to the flexibility of phenylalanine, we find that BVM preferentially coordinates to a subset of phenyl groups; in our simulations, we see these phenyl groups flatten, thereby establishing contacts with the pentacyclic moiety, while the remaining phenyl groups extend into the vacated space within the central pore and the 6HB distorts accordingly. To improve BVM coordination, these key residues likely need to be mutated to hydrophobic residues with stiff side chains, such as branched amino acids. For instance, M367I has been shown to have high BVM efficacy.44

Second, escape mutations can disrupt the hydrophobic knob-in-hole (KIH) interfaces16 that are essential to the quaternary structure of the 6HB (orange box in Figure 6). Here, the “knobs” consist of methyl side chains from A360 and A364 that coordinate within “holes” that are formed between side chains from the adjacent helix (the next counterclockwise helix in the 6HB in Figure 6). These side chains are composed of V362, L363, and V366 for A360 and A364, and M367, and V370 for A364. The asymmetry between the knob and hole side chains is likely important as the smaller methyl knobs can be coordinated by the bulkier side chains that form the hole. Hence, increasing the knob size with A364V mutants or decreasing the hole size with V370A mutants leads to loss of...
KIH packing efficacy, resulting in loss of 6HB symmetry. In these cases, we find that BVM interactions with the interior hydrophobic interface of the 6HB (cooperatively enhanced when IP6 is present) are insufficient to retain 6HB symmetry. Although we did not simulate it, mutants may instead stabilize the 6HB such that BVM has negligible perceived efficacy. In this case, mutant virus in the absence of BVM can sufficiently symmetrize the 6HB to prevent protease access. On the basis of the aforementioned KIH interactions, one way to do so may be to mutate hole-forming residues with branched (i.e., bulkier and stiffer) and more hydrophobic side chains, e.g., valine, leucine, and isoleucine. Indeed, L363I, A366V, and M367I mutants have been shown to decrease viral infectivity and CA/SP1 processing, while BVM itself had indiscernible effects on L363I mutant virus.\textsuperscript{14,44} The T371I mutant is also known to prevent maturation and phenocopies BVM,\textsuperscript{15,45} while BVM likely induces no additional symmetry preservation in the 6HB.

Our findings highlight difficult constraints that must be overcome by next-generation maturation inhibitors, which include precise coordination to the 6HB in both WT and polymorphic viruses. The dimethylsuccinyl moiety has been recognized as a pharmacophore\textsuperscript{32,46} for BVM, and our analysis shows that its coordination to K359 has potent 6HB symmetry preservation effects. Interestingly, this effect is cooperatively enhanced by the presence of IP6, which suggests modification of the dimethylsuccinyl moiety to improve overall K359 coordination may be valuable. The recognition that the QVT motif is a major source of viral polymorphism has motivated numerous modifications near the C-28 carboxylic acid moiety,\textsuperscript{30,32,33,35,47,48} which have demonstrated that it is possible to overcome loss of susceptibility due to polymorphism. On the basis of our results, we suggest that modification or replacement of the pentacyclic triterpenoid moiety is a potential third avenue. As this moiety establishes key contacts with the hydrophobic interior of the 6HB, it may be possible to emulate the effect of mutants that simultaneously stabilize KIH and inhibitor interactions. For instance, a triterpenoid with larger hydrophobic (e.g., propyl) groups near the terminal cyclic rings may aid in M367 interactions and facilitate downstream KIH interactions. Finally, this work suggests the need to evaluate inhibitor activity from the perspective of structural ensembles, and to that end, we anticipate that molecular simulations will be a valuable tool for fundamental inhibitor studies and design. The dynamical mechanisms of protease access to the 6HB remain an open fundamental question, and it would be interesting to investigate whether particular states in the 6HB structural ensemble facilitate proteolytic cleavage.

**CONCLUSIONS**

Using extensive MD simulations and free energy sampling, we have provided a comprehensive comparison of molecular structure–activity relationships across WT HIV-1 and mutant (L363F, A366V, and V370A) CA\textsubscript{CTD}/SP1 viruses in the presence and absence of BVM, a maturation inhibitor. Our analysis reveals an ensemble of 6HB structures with varying degrees of 6-fold symmetry. The most important conclusion is that the inclusion of BVM in WT virus rigidifies the 6HB through multiprotomer interactions, thereby preserving 6-fold symmetry, which is cooperatively enhanced by IP6, an assembly cofactor. The escape mutants reduce 6-fold symmetry throughout the 6HB, which is partially recovered by BVM in particular cases. Our free energy calculations demonstrate that loss of 6-fold symmetry facilitates helical uncoiling, which precedes proteolytic cleavage.\textsuperscript{19,22} Consequently, the efficacy of BVM is due to additional resistance to symmetry loss. Two primary escape mechanisms were identified: (i) failure to mitigate lost knob-in-hole 6HB interactions and (ii) failure to uniformly coordinate the 6HB within the central pore. Our analysis demonstrates the utility of MD simulations to provide insight into the molecular basis for the stabilization of the CA\textsubscript{CTD}/SP1 6HB conferred by BVM, which provides new directions for the design of future HIV maturation inhibitors. In particular, our findings imply that maturation inhibitor design should focus on symmetry-inducing interactions within the multiprotomer bundle rather than those that impart high affinity.

**METHODS**

**Molecular Dynamics Simulations.** All MD simulations were prepared using hexameric CA\textsubscript{CTD}/SP1 domains of an atomic model (PDB code 5L93\textsuperscript{37}), which was modified accordingly with the point mutations L363F, A366V, and V370A. Systems prepared with IP6 and BVM were initialized with Monte Carlo insertion of the molecule in the central pore of the hexamer near K359 and M367, respectively. Each system was solvated by water in a dodecahedral domain with at least 1.6 nm of space between the edges of the protein and the domain. Water molecules were then randomly replaced with 0.15 M NaCl. The total size of each system was ~108 000 atoms with initial box lengths of 11.6 nm × 11.6 nm × 8.2 nm.

All simulations used the CHARMM36m force field\textsuperscript{40} and were performed using GROMACS 2016.\textsuperscript{50} Minimization was performed using steepest descent until the maximum force was reduced to 1000 (kJ/mol)/nm. Then, equilibration was performed in several phases. First, 10 ns was integrated in the constant NVT ensemble using the stochastic velocity rescaling thermostat\textsuperscript{51} with a damping time of 0.2 ps and a time step of 1 fs. During this phase, the Cβ backbone of the protein was harmonically restrained with a force constant of 1000 kJ/mol/nm\textsuperscript{2}. An additional 40 ns was integrated in the constant NPT ensemble using the stochastic velocity rescaling thermostat\textsuperscript{51} (2 ps damping time) and the Parrinello–Rahman barostat\textsuperscript{52} (10 ps damping time) and a time step of 2 fs. Finally, an additional 750 ns was integrated in the constant NVT ensemble using the Nose–Hoover chain thermostat\textsuperscript{53} (2 ps damping time) and a time step of 2 fs. During the last three phases, the Cβ backbone of helix 9 throughout the hexamer was harmonically restrained with a force constant of 500 kJ/mol/nm\textsuperscript{2}. Throughout this procedure, H-containing bonds were constrained using the LINCS algorithm.\textsuperscript{54} Statistics were gathered every 200 ps, and the final 600 ns of each trajectory was used for analysis, which was performed using a combination of VMD Tcl scripts\textsuperscript{55} and Python scripts using MSMBuilder\textsuperscript{56} and MDTraj.\textsuperscript{57}

Free-energy calculations were performed using PLUMED 2.4\textsuperscript{58} and GROMACS 2016.\textsuperscript{68} Final configurations from the MD simulations for each of the three reported systems were used as initial configurations. For each system, five independent replicas were prepared using random velocity distributions and equilibrated in the constant NVT ensemble (same thermostat and time step as above) for 200 ns; the same restraints on helix 9 and H-containing bonds were kept throughout these simulations. Well-tempered metadynamics\textsuperscript{56} was subsequently performed for 4.6 μs by biasing the AB\textsubscript{eq} collective variable for a single helix; Gaussian hills with 0.5 kJ/mol height and 0.05 width were deposited every 1 ps with a bias factor of 15 kJ/T. Data were collected every 1 ps, and the Tiwary–Parrinello algorithm\textsuperscript{69} was used to reweight FESs onto a 2D projection along AB\textsubscript{eq} and y\textsubscript{eq}. Here, y\textsubscript{eq} was computed using the nonbiased helices, i.e., only four (out of six) angles were computed and averaged, as the biased helix twists out of the 6HB when unfolding. All reported FESs were averaged across replicas after aligning free energy values at the observed global minimum.
Details of Structural Analysis. We computed gyration moments on a per-ring basis along the 6HB (see Figure 1). For each residue with index $i$ between H336 and T371, we calculated the mean ($\mu_i$) of the distance between each of the six C atoms of residue $i$ ($r_{C_i}$) and their collective center of mass ($r_{COM}$):

$$\mu_i = \frac{1}{6} \sum_{k=1}^{6} (r_{C_{ik}} - r_{COM})$$  \hspace{1cm} (1)

We then calculated the second through fourth moments about $\mu_i$, i.e., $\mu_{ij}$ through $\mu_{ijk}$:

$$\mu_{ij} = \frac{1}{6} \sum_{k=1}^{6} (r_{C_{ik}} - r_{COM})^2$$

$$\mu_{iij} = \frac{1}{6} \sum_{k=1}^{6} (r_{C_{ik}} - r_{COM})^3$$

$$\mu_{iijj} = \frac{1}{6} \sum_{k=1}^{6} (r_{C_{ik}} - r_{COM})^4$$  \hspace{1cm} (2)

Therefore, for each 6HB conformation, a total of 64 features were collected. Our use of features that represent the collective configuration of the six helices ensures permutation invariance across helix labels. The features from all 120 trajectories (i.e., across all replicas for WT and mutant virus) were used for dimensional reduction. The $\psi_{6,2D}$ for each $C_\alpha$ ring along the 6HB was quantified as follows:

$$\psi_{6,2D} = \frac{1}{10} \sum_{j=369}^{370} \sum_{k=1}^{6} \exp(i\theta_{jk})$$

where $\theta_{jk}$ is the angle formed between the $C_\alpha$ atom of residue $j$ in helix $k$, the center of mass (COM) of all $C_\alpha$ atoms of residue $j$, and the $C_\alpha$ atom of residue $j$ in helix $k + 1$.

The $\omega_{sim}$ was computed using the following:

$$\omega_{sim} = \frac{1}{6} \sum_{k=1}^{6} \frac{1}{14} \sum_{i=357}^{361} \frac{1}{2} (1 + \cos(\theta_{ik} - \theta_{jk}))$$

where $\theta_{ik}$ is the angle formed between the $C_\alpha$ of residue $i = 1, 6$ and $i + 1$ in helix $k$ and $\theta_{jk}$ is the expected angle for $\alpha$-helices ($\approx 91^\circ$).

The $\alpha_{sim}$ was calculated using the following:

$$\alpha_{sim} = \frac{1}{6} \sum_{k=1}^{6} \frac{1}{14} \sum_{i=357}^{361} \frac{1}{2} (1 + \cos(\psi_{i} - \phi_{ij}))$$

where $\phi_{ij}$ ($\psi_{i}$) is the N–C$_\alpha$ (C$_\alpha$–C) backbone torsional angle of residue $i$, and $\phi_0$ ($\psi_0$) is the reference angle ($-60^\circ$ for both) for $\alpha$-helices; we focus on residues 361–367 as the scissile bond is located between residues 363 and 364.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.1c08922.

Six additional figures showing (i) time-series profiles of positions for BVM, IP6, and $C_\alpha$ within the 6HB, (ii) RMSD comparison between averaged MD and experimental 6HB structures, (iii) $\alpha$-helix similarity of the $C_\alpha$ backbone ($\omega_{sim}$) throughout the 6HB for each cluster, (iv) a representative schematic of BVM and IP6 coordinating to residue K359 within the 6HB, (v) time-series profiles of BVM orientation within the 6HB, and (vi) proximity contact maps between BVM and residues 363/367 for clusters 3, 6, and 7 and a table summarizing the statistics depicted in Figure 3 (PDF).

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Notes

The authors declare no competing financial interest.

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