Off-Pathway Assembly: A Broad-Spectrum Mechanism of Action for Drugs That Undermine Controlled HIV-1 Viral Capsid Formation

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Supporting Information

ABSTRACT: The early and late stages of human immunodeficiency virus (HIV) replication are orchestrated by the capsid (CA) protein, which self-assembles into a conical protein shell during viral maturation. Small molecule drugs known as capsid inhibitors (CIs) impede the highly regulated activity of CA. Intriguingly, a few CIs, such as PF-3450074 (PF74) and GS-CA1, exhibit effects at multiple stages of the viral lifecycle at effective concentrations in the pM to nM regimes, while the majority of CIs target a single stage of the viral lifecycle and are effective at nM to μM concentrations. In this work, we use coarse-grained molecular dynamics simulations to elucidate the molecular mechanisms that enable CIs to have such curious broad-spectrum activity. Our quantitatively analyzed findings show that CIs can have a profound impact on the hierarchical self-assembly of CA by perturbing populations of small CA oligomers. The self-assembly process is accelerated by the emergence of alternative assembly pathways that favor the rapid incorporation of CA pentamers, and leads to increased structural pleomorphism in mature capsids. Two relevant phenotypes are observed: (1) eccentric capsid formation that may fail to encase the viral genome and (2) rapid disassembly of the capsid, which express at late and early stages of infection, respectively. Finally, our study emphasizes the importance of adopting a dynamical perspective on inhibitory mechanisms and provides a basis for the design of future therapeutics that are effective at low stoichiometric ratios of drug to protein.

INTRODUCTION

Over the past few decades, antiretroviral therapy (ART) for human immunodeficiency virus type 1 (HIV-1) has made substantial progress. These advances can be attributed, in part, to the identification of multiple enzymes that are critical to the HIV-1 lifecycle, such as reverse transcriptase, integrase, and protease, as well as the development of small molecules that competitively inhibit their activities. Current state-of-the-art ARTs consist of a combination (cART) approach that utilizes multiple drugs, each active against different targets; this strategy is also referred to as highly active ART (HAART). Nonetheless, the primary challenges for HAARTs are to maintain efficacy, safety, and tolerability. In particular, genetic variation is responsible for drug resistance in patients. Treatment regimens must therefore adapt to these circumstances, e.g., through the introduction of drugs with alternative mechanisms of action, since clinically available options are limited.

An emerging class of drugs aims to disrupt the activity of the Gag (group-specific antigen) polyprotein, which is responsible for coordinating the late stages of the viral lifecycle. The capsid domain (CA) of Gag is an attractive therapeutic target, since both the assembly and maturation of infectious viral particles are mediated by interactions between CA. During maturation, for example, CA self-assembles into a conical capsid (i.e., the mature core), composed of more than 1000 CA monomers, that encases the viral genome. Given the functional significance of the CA, it is also important to note that its sequence is highly conserved (around 70%) among HIV-1 subtypes, thereby reducing the risk of viral polymorphism. Drugs that target CA are known as capsid inhibitors (CIs) and have been studied for nearly a decade. Several candidates have been identified that demonstrate the feasibility of the CI approach, such as Bevirimat and PF-3450074 (PF74), which have half maximal effective concentrations (EC50), a measure of drug potencies, at nM to μM concentrations. Most recently, GS-CA1 was introduced as a promising CI with an EC50 around 85 pM concentration, and targets the same binding pocket as PF74 during early and late stages of viral infection. Nonetheless, a clinically viable CI has yet to be discovered. A molecular understanding of the mechanism of action for CIs is currently lacking, and represents a barrier for the development of new therapeutics.

Perhaps the most extensively studied CI, PF74 serves as a useful example to highlight the complexity of potential mechanisms for CIs. Initial reports of PF74 have suggested that capsid destabilization is the primary mechanism of action for the drug. Further capsid disassembly assays have shown that PF74 induces viral uncoating, although a contradictory study, using assays at similar concentrations of PF74, observed no discernible effects on viral uncoating and reverse
transcription.29 Interestingly, similar assays that were performed on pre-assembled CA tubules have shown that PF74 has a stabilization effect.28,30,31 Several binding assays have indicated that PF74 preferentially binds to CA multimers rather than isolated CA23,28,31,32 and have found that CA assembly rates increase with PF74 present.23,31 One hypothesis that may resolve the aforementioned contradiction is that PF74 affects the kinetics of the CA assembly process, which warrants further investigation. Another hypothesis has also been presented on the basis of the crystal structure of PF74 in complex with CA hexamers. Since the drug binds to a pocket at the interface between the N-terminal domain (NTD) and C-terminal domain (CTD) of adjacent CA monomers,23,28 PF74 binding may preclude variable curvatures in CA oligomers that must be adopted for closed capsids. However, since this binding pocket has also been associated with cleavage and polyadenylation specific factor 6 (CPSF6) and nucleoporin 153 kDa (NUP153), two cellular transport factors that aid nuclear import,33,34 a third possibility is that nuclear integration of the virus is abrogated due to competitive inhibition by PF74.29 Taken together, these observations suggest that PF74 (and related CIs, such as GS-CA1) can exhibit complex, multimodal mechanisms of action that target viral maturation, uncoating, reverse transcription, and nuclear import. Understanding the molecular mechanisms by which CIs produce one key aspect (capsid assembly/disassembly) of such broad activity is the focus of this work.

We use coarse-grained (CG) molecular dynamics simulations based on prior CG models11 to investigate the impact of CIs on viral infectivity. We hypothesize that the role of CIs is to perturb the population of small, intermediate oligomers of CA. These effects are implicitly incorporated into our CG model and used to examine the subsequent implications on late and early stages of the viral lifecycle. We simulate both capsid assembly and disassembly and emphasize the differences between the assembly/disassembly pathways in models for both holo and apo capsids. Our findings suggest that CIs have multiple modes of action by increasing the population of mature capsids that either (i) fail to enclose viral RNA during maturation or (ii) spontaneously disassemble before transport and nuclear integration. Furthermore, our study implies that the targeting of CA oligomers, which requires low stoichiometric loadings of drugs to CA, is a possible design strategy for future therapeutics.

## RESULTS AND DISCUSSION

We begin by considering the hierarchical nature of viral capsid assembly, which is schematically shown in Figure 1. The capsid is an enclosed protein core that is composed of tiled CA hexamers with 12 CA pentamers incorporated for topological completeness.17,18 In solution, CA exists in a dynamic equilibrium between CA monomers and dimers.35,36 Furthermore, conformations of CA dimers are inherently dynamic, as depicted in the top row of Figure 1, and importantly, only adopt conformations compatible with the mature capsid with S~10% probability (i.e., the conformer observed in mature CA hexamers, which is shown in the upper half of the red box in Figure 1).37 CA dimers associate during the assembly process to form the final capsid structure. Prior CG simulations10,11,38 and a kinetic model39 have shown that a key intermediate is the trimer of dimers (TOD) structure, depicted in the lower half of the red box in Figure 1. Three TODs form a complete CA hexamer that can simultaneously template adjacent hexamers. As CA oligomers increase in size, there are numerous possible assembly outcomes, including stalled or malformed assemblies. However, under precise conditions of weak CA-CA interactions with highly specific association interfaces, which are only accessible in a limited manner, the self-assembly process expresses the characteristic fullerene core morphology.71 These results have suggested that the nucleation of specific intermediate oligomers is an essential
aspect of canonical mature core assembly. It is also notable that self-assembly is predicted to proceed even in the absence of viral RNA, which is consistent with recent experiments that have demonstrated mature capsid formation when co-assembled with inositol phosphate, a small-molecule co-factor. The conditions used in previous simulations implicitly include the putative effects of these co-factors, and the simulations presented hereafter similarly assume that RNA is not essential for core formation.

To consider the impact of CIs on the CA assembly process, we note that crystallographic structures show evidence that PF74 and GS-CA1 preferentially bind to an inter-CA pocket between the respective NTD and CTD domains of two adjacent CA monomers. Importantly, these pockets only appear when CA oligomerizes, such as in TODs, and appear to mediate inter-CA interactions, thereby stabilizing multimeric CA. We therefore hypothesize that CIs stabilize the interface between CA domains, increasing the population and/or lifetime of intermediate oligomers. A natural question to investigate is how these perturbations affect the overall CA assembly process.

To simulate the influence of CIs, we use the “ultra-CG” (UCG) model\(^\text{11}\) that was previously developed to investigate mature capsid assembly. Briefly, the UCG model consists of a \(\alpha\) resolution representation of CA helices with an elastic network model\(^\text{13}\) (ENM) used to conserve the secondary and tertiary structure of the protein. Select inter-CA contacts that are exposed in the “mature” configuration are identified from X-ray structures\(^\text{19,14}\) and projected as “virtual” CG sites with two possible states: (1) a non-interacting “inactive” state and (2) an interacting (through an attractive Gaussian interaction) “active” state. Throughout the simulation, CA dimers periodically switch between active (\([\text{CA}^+]\)) and inactive (\([\text{CA}^-]\)) states and a constant ratio between the two is maintained; see Figure S1 for more details. Here, we include the effect of CI binding to CA by introducing a fixed population of \([\text{CA}^+]\) that have pre-assembled into TODs (\([\text{CA}^+]\)\(_{\text{TOD}}\)), as highlighted by the red box in Figure 1. The TODs are maintained by an auxiliary ENM that conserves the relative position of adjacent dimer domains as depicted in Figure S1. We note that the additional ENM also rigidifies the TOD and restricts the distribution of curvatures adopted by the oligomer, which is consistent with all-atom simulations that we analyze in Figure S2. We performed CG MD simulations under conditions that previously resulted in self-regulated assembly\(^\text{13}\) (\([\text{CA}^+]\)/\([\text{CA}^-]\)  = 0.11, \([\text{CA}]\) = 4 mM, inert crowder density at 200 mg/mL, UCG state switching interval at \(5 \times 10^5\) timesteps) with \([\text{CA}^+]\)\(_{\text{TOD}}\)  = 0.025, 0.050, 0.075, 0.100, 0.200, and 0.500 mM and 2 replicates for each condition resulting in a total of 12 independent trajectories; note that \([\text{CA}^+]\)\(_{\text{TOD}}\)  = 0.025 mM represents a single TOD out of a total of 616 CA dimers in a 80 nm length cubic box. Each simulation was run for \(2 \times 10^8\) timesteps \((\tau)\) with \(\tau = 10\) fs in CG time (which is not to be confused with actual time). Further details can be found in Methods and in ref. 11.

The bottom row in Figure 1 depicts the outcomes of each of the 12 CG simulations. Here, we qualitatively define three classes of core morphologies for the purposes of the following discussion: (1) canonical, (2) eccentric, and (3) malformed. Malformed capsids are those with non-contiguous, semi-amorphous lattices, such as when multiple aggregates of CA merge. Canonical and eccentric capsids refer to contiguous lattices that are distinguished by their curvatures, and relatedly, their pentamer distributions; in the former case, pentamers are dispersed such that all pentamers are separated by (at minimum) a hexamer, while the latter case contains adjacent pentamers in pentamer-dense regions, which also denote regions of high curvature; the need for this distinction will become evident during the discussion below. From our CG simulations, we observe 1 canonical, 10 eccentric, and 1 malformed capsid structure. We note that an explicit trend between final core morphology and \([\text{CA}^+]\)\(_{\text{TOD}}\) as summarized in Table 1, is not observed at present, although this can be attributed to the stochastic nature of the process; additional independent sampling at each \([\text{CA}^+]\)\(_{\text{TOD}}\) would be required to confirm a potential trend with respect to \([\text{CA}^+]\)\(_{\text{TOD}}\) but is currently computationally cost prohibitive and outside the scope of this work. Nonetheless, insight into the influence of \([\text{CA}^+]\)\(_{\text{TOD}}\) on structural aspects of the assembly process can be obtained, and certainly with respect to inhibitor-free systems as discussed later. Our simulations relatedly suggest that the majority of cores that form due to CIs are highly pleomorphic with many cores exhibiting large degrees of curvature. This is a notable result when compared to the expected behavior of wild-type (WT) viruses. Previous cryo-electron microscopy (cryo-EM) surveys of mature WT virions have revealed a variety of capsid phenotypes, including conical capsids, tubular capsids, incompletely capsids and dual capsids, although the majority (around 60 to 90%) appears to be conical. Our simulations suggest that the presence of CIs shifts this distribution toward the formation of non-canonical cores, which is consistent with cryo-EM images of virions treated by GS-CA1. The wide distribution of end-point morphologies shows that even slight positive perturbation to TOD populations has profound effects on CA assembly that favor alternative pathways. We note that this view is conceptually consistent with recent reports of pathway complexity observed in supramolecular polymers, which emerges when assembly kinetics cannot be described by simple nucleation-and-elongation models.\(^\text{47-49}\) We next discuss the origins of the distinct pleomorphism in lattice morphology that emerges due to the presence of CIs.

To characterize the lattice morphology, we first recognize that the CA lattice can be represented as a connected graph. One may conceive of each CA monomer as a vertex \((v)\) on a graph \((G)\) with an edge \((e)\) between vertices signifying their proximal nature. Within the assembled CA lattice, \(e\) represents dimeric CA as well as adjacent intra-capsomer CA; in other words, each \(v\) is ideally connected to three neighboring \(v\) within \(G\), which strictly represents the largest assembled CA lattice. Throughout the simulation trajectories, we use a distance criterion to construct \(G\):
Figure 2. Topology of the assembled CA lattice is assessed by the eccentricity ($\lambda$) of each vertex (i.e., each CA monomer) in a graph representation of the lattice. We show the distribution of $\lambda$ ($P(\lambda)$) at select points during one trajectory of $[CA^+]_{CI} = 0.100$ mM for demonstrative purposes: at (a) $1 \times 10^8$, (b) $7 \times 10^8$, and (c) $14 \times 10^8$ MD timesteps. The same analysis is performed in (d) using half of the lattice from (c), i.e., after cleavage by a plane perpendicular to the long axis of the capsid. Representative lattices are depicted as insets in (a-d) with each monomer shown as a sphere colored by its $\lambda$ from red ($\lambda = 0.5$) to blue ($\lambda = 1.0$). The topology of the lattice can be qualitatively characterized by the skew of the distribution of $\lambda$ ($\gamma(\lambda)$) in which (a,d) zero skew is indicative of an open lattice with isotropic edges, (b) negative skew is indicative of an open anisotropic lattice, and (c) positive skew is indicative of a closed, non-spherical lattice.

Figure 3. Assembly time-series plots that depict (a) the number of assembled hexamers, (b) the number of assembled pentamers, (c) the number of incorporated inhibitor-bound TODs (TODCI), and the (d) skew ($\gamma(\lambda)$) and (e) kurtosis ($\kappa(\lambda)$) of the distribution of eccentricities ($\lambda$) throughout the assembled lattice as a function of CG MD time step (shifted with respect to the onset of lattice growth) for each system within the listed concentration of capsid inhibitors (CIs). We find accelerated assembly, especially with respect to pentamers, in the CI-present simulations, which appear to be commensurate with negative $\gamma(\lambda)$ and $\kappa(\lambda)$, i.e., a descriptor that indicates anisotropic edge growth in the protein lattice. The shaded region depicts the standard deviation around the mean from four trajectories within the listed range of concentrations while the solid line depicts a single trajectory. In (f), molecular snapshots of the protein lattice for the 0.100 mM case (green line in (a)−(e)) at the listed MD time step ($\tau [\times 10^5]$) are shown; green, red, and blue NTD domains in capsomers indicate hexamers, pentamers, and incomplete capsomers, respectively.

\[ G = \{V, E\} \]  
(1) \[ V = \{v|v \in \text{CA lattice}\} \]
$$E = \{e_{ij} = (v_i, v_j) \mid d_{NTD}(i, j) < r_c \text{ or } d_{CTD}(i, j) < r_c\}$$  

(3)

with $V$ and $E$ denoting the set of $v$ and $e$ in $G$, $i$ and $j$ denoting the index of each CA monomer, $d_{NTD}$ ($d_{CTD}$) denoting the measured distance between residue V36 (E180) in $i$ and $j$, and $r_c$ denoting a cutoff distance of 2.75 nm; further justification for this cutoff distance and choice of residues is included in Figure S3. Next, we consider a property of graphs known as eccentricity ($ecc$), which effectively measures how "far" a given vertex is from its furthest vertex. The $ecc$ of each $v$ is formally defined as

$$ecc(v) = \max(\minpath(v, v) \mid i \neq j; v, v_j \in G)$$

(4)

which quantifies the maximum path distance among the set of minimum paths between $v_i$ and every other $v$ in $V$. For our purposes, we consider the normalized $ecc$ ($\lambda = ecc(v)/\max\{ecc(v)\}$) and statistical measures on its distribution $(P(\lambda))$ as a metric to classify CA lattice morphologies.

The utility of $P(\lambda)$ can be shown with four representative CA lattices (extracted from a single trajectory) that are analyzed in Figure 2; additional discussion on $P(\lambda)$ can be found in Figure S4 and Table S1. Figure 2a depicts an early stage of CA nucleation in which the lattice appears largely two-dimensional and isotropic around its six-fold (C6) axis. The resultant $P(\lambda)$ exhibits a normal distribution with the smallest (largest) $\lambda$ associated with the center (edges) of the lattice. In Figure 2b, the lattice has grown with some edges appearing "jagged" (a nascent dendrite, in some sense), which is indicative of anisotropic nucleation. The $\lambda$ associated with $v$ that participate in these jagged edges is large (e.g., $\lambda > 0.8$) in comparison to interior $v_i$ in fact, as these edges become more dendritic, we expect large $\lambda_i$ to be increasingly favored in $P(\lambda)$. Here, we argue that anisotropic growth at the edges of the CA lattice are relevant since the branching junction between dendrites may nucleate into pentamers, presumably to anneal the lattice and to minimize potential strain due to curvature. In Figure 2c, the lattice has fully wrapped into an enclosed 2D lattice, such that all $v$ can be considered interior $v$. Now, $P(\lambda)$ favors a higher population of small $\lambda_i$ (e.g., $\lambda_i < 0.8$) with large $\lambda_i$ associated with oblong areas of the capsid; note that pristine cleavage of the capsid, such that the edges are largely composed of hexamers that are a uniform distance from the narrow end of the capsid, recovers a $P(\lambda)$ with a normal distribution, as seen in Figure 2d. This analysis (and the additional discussion in Supporting Information) suggests that we may use the skew and kurtosis of $P(\lambda)$ (i.e., $\gamma(\lambda)$ and $\kappa(\lambda)$) to qualitatively classify CA lattice growth; $\gamma(\lambda) \approx 0$ and $\kappa(\lambda) < 0$ signifies a lattice with edges that are isotropically distant from the lattice center while $\gamma(\lambda) < 0$ and $\kappa(\lambda) < 0$ (or $\gamma(\lambda) > 0$ and $\kappa(\lambda) > 0$) represents a lattice with anisotropic edges (or with contiguous and uniform oblong character, such as in complete enclosures).

On the basis of the aforementioned metric, we now analyze time-series profiles in Figure 3, in which we compare CG trajectories from CI-bound cases (12 trajectories) to that of non-CI cases (four trajectories); two of the non-CI trajectories were obtained from previous work while two additional trajectories were generated using the same conditions. We find that in all cases (both CI-bound and non-CI) the assembly rate of CA hexamers (Figure 3a) is comparable with the exception of $[CA^+]_{CI} = 0.200–0.500$ mM, which is accelerated by nearly a factor of 2 compared to the $[CA^+]_{CI} = 0.000$ mM case. More evident differences are observed in the assembly rate for pentamers (Figure 3b). Here, pentamer incorporation rates are positively correlated with an increase of $[CA^+]_{CI}$. The observed increase in multimerization rate is consistent with experimental multimerization assays for PF74. Our simulations suggest that CI-bound populations of CA (even at stoichiometric ratios as low as 1 mol%) can have a profound impact on multimerization rates. Interestingly, the onset of pentamer incorporation is notably earlier in the CI-bound cases (around $50–100 \times 10^6$) compared to the non-CI case (around $250–350 \times 10^6$). The incorporation of CI-bound TODs (Figure 3c) is observed to directly precede pentamer incorporation and suggests that CIs influence core morphology by promoting pentameric defects; the origins of this behavior will be discussed below.

Analysis of the changes to the topological character of the growing CA lattice may provide further insight into the effect of CIs on CA nucleation and growth. In all cases (CI-bound and non-CI), $\gamma(\lambda)$ and $\kappa(\lambda)$, as seen in Figures 3d,e, is initially close to 0.0 and −1.0. The former, however, shifts toward negative values directly preceding the onset of pentamer incorporation (and commensurate with CI-bound TOD incorporation in the CI-bound cases); recall that the combination of negative $\gamma(\lambda)$ and negative $\kappa(\lambda)$ is a feature indicative of non-uniform protrusions along the lattice edges. The resultant pentamer incorporation typically seems to anneal the lattice such that $\gamma(\lambda)$ increases toward values greater than or equal to 0.0, and the rate of pentamer incorporation is temporarily suppressed. However, as the edges of the lattice continue to nucleate CA, anisotropic character may re-emerge and promote the additional incorporation of pentamers. The edges of the lattice may ultimately anneal to form an enclosed CA lattice (with $\gamma(\lambda)$ and $\kappa(\lambda) \geq 0$). Here, CA within incomplete capsomers, which are likely to be in areas of high curvature, continue to relax until pentamers are formed, e.g., as evident by the time-series profiles in which pentamer counts increase while hexamer growth is stalled. Molecular snapshots from a CI-bound case that exhibits all of these steps are shown in Figure 3f.

We interpret these collective results by proposing that CI-bound TODs (i.e., the fixed TODs in our simulations) accelerate CA assembly compared to CI-free cases by increasing the number of accessible assembly pathways. First, let us consider that TODs, while important building blocks, still require free CA dimers (and potentially, other small oligomers) to continue assembly of the CA lattice. Each CA dimer favors two binding interfaces that should be satisfied for successful association, as seen in Figure 4a. Hence, continued CA assembly requires the near-simultaneous fastening of two CA dimers at adjacent edges of the lattice, which subsequently create new edges for future second-order CA association events. The increased population of TODs, on the other hand, enables CA lattice growth through the near-simultaneous binding of a TOD and CA dimer. In this alternative assembly pathway, new binding edges also appear. It is instead likely that subsequent assembly steps only require a single CA dimer, as seen in Figure 4a, which therefore accelerates CA assembly. We note that at low $[CA^+]_{CI}$ (e.g., the 0.025–0.050 mM regime in this work) this alternative assembly pathway is only feasible when the initial nucleation of the CA lattice is independent of the CI-bound TOD. In the event that the CI-bound TOD is the initial nucleator and the reservoir of available CI-bound TODs is depleted, the canonical assembly pathway is expected to proceed; this scenario is consistent with...
the $[\text{CA}^+]_{\text{Cl}} = 0.050$ mM case that resulted in a canonical core (see Table 1).

To demonstrate the emergence of additional assembly paths in the presence of CIs, we perform 50 short independent simulations (over $150 \times 10^6$ $\tau$) of CA assembly with $[\text{CA}^+]_{\text{Cl}} = 0.000$ and 0.100 mM starting from a pre-assembled lattice with 37 CA hexamers (i.e., the equivalent of four concentric “layers” of hexamers). We depict the progression of $\gamma(\lambda)$ in Figure 4b,c and compare the two cases; we focus on $\gamma(\lambda)$ as $\kappa(\lambda)$ remains consistently negative during this early growth phase (see Figure 3). In the $[\text{CA}^+]_{\text{Cl}} = 0.000$ mM case (Figure 4b), we find that the majority of trajectories maintain $\gamma(\lambda)$ around 0.0. However, in the $[\text{CA}^+]_{\text{Cl}} = 0.100$ mM case (Figure 4c), a fraction of the trajectories instead adopt $\gamma(\lambda) < 0$, which we suggest is commensurate with increasingly anisotropic lattice growth, while this fraction increases as time progresses. This observation is most evident after $50 \times 10^6$ $\tau$ states associated with $\gamma(\lambda) < -0.75$ are sampled by CI-bound trajectories (see red box in Figure 4c) while absent from non-CI trajectories (see red box in Figure 4b). We therefore suggest that the presence of CI-bound TODs enables an alternative assembly pathway through enhanced CA association kinetics and increasingly anisotropic lattice growth. Here, we should mention that our UCG model does not explicitly consider the possible binding of CIs to CA after multimerization. However, as we expect CIs to stabilize CA association, thereby accelerating nucleation and growth (or hindering CA reconstruction), the qualitative impact on assembly is likely comparable to that of the current UCG model.

One consequence of the accelerated CA assembly, and relatively, the rapid pentamer incorporation, induced by CIs is that the resultant mature cores may fail to enclose viral RNA for two reasons. First, the enhanced CA assembly kinetics may preclude the need for viral RNA and nucleocapsid (NC), i.e., the ribonucleoprotein (RNP) complex, to serve as templates for assembly; some experiments have suggested that the RNP complex nucleates mature CA assembly, which is further mediated by integrase, although this behavior does not appear to be universal as evident by mature capsid formation even in the absence of an enclosed RNP complex. Second, CI-perturbed cores are likely to adopt highly curved morphologies with small internal volumes that may impede the encapsulation of the RNP complex. In our simulations, cores that underwent full enclosure were observed to contain on the order of 100 hexamers (see Figure 3a). For comparison, previous accounts from cryo-EM data suggest canonical capsid cores contain on the order of 200 hexamers. Based on an effective reduction of surface area by a factor of 2, we approximate an associated reduction of internal volume by up to 65% (for simple spherical geometries). However, we note that RNA condensation, which is likely dependent on salt and viral protein content, may allow the RNP complex to occupy these reduced volumes; note that such condensation is suggested by the localization of RNP-associated densities in cryo-EM images within the broad region of conical mature capsids. Nonetheless, the proposed phenotype is consistent with previous cryo-EM experiments that have observed RNP-associated density outside that of protein cores, especially those treated with inhibitors. Hence, our simulations suggest that one mechanism of action of CIs is to increase the population of eccentric cores that may fail to enclose the RNP complex.

In the event that CI-bound mature cores successfully condense the RNP complex, we also investigated their viral uncoating behavior. During the post-entry stages of viral infection, capsid disassembly and trafficking toward the nuclear pore is highly regulated. Although the uncoating process is not completely understood, it has been suggested that a combination of biological triggers, such as the binding of host cell factors or the initiation of reverse transcription, act as signals to delay then initiate disassembly. For instance, cyclophilin A (cypA) can stabilize the mature core throughout trafficking and disengage in a downstream event to allow induced uncoating by transportin 3 (TNPO3). On the other hand, restriction factors (such as TRIM5α and TRIMCyp) can induce premature uncoating. Maintenance of the viral capsid post-entry is additionally purported to be an essential barrier against innate immune responses. Hence, perturbations to canonical uncoating behavior that result in spontaneous uncoating may also reduce infectivity.

We simulate putative post-entry events following the procedures established in ref 11, in which we deplete the
concentration of available [CA\(^+\)] (= 0 mM) and integrate dynamics over 12 \times 10^8 \tau. In Figure 5, we compare time-series profiles that depict the fraction of the mature core that persists. Here, we select and compare four closed end-point morphologies from our simulations described above. As a control, we also prepared a closed canonical core that was pre-assembled based on alignment to an atomic model for a complete capsid (PDB 3J3Y). As seen in Figure 5, the 3J3Y core remained stable throughout our simulated time-scale, in agreement with previous all-atom MD simulations that suggest the core is inherently stable at equilibrium.

Interestingly, three out of four of the selected eccentric cores undergo a clear stepwise disassembly process, as described by the processive periods of plateaus and decrements seen in Figure 5a. The molecular snapshots depicted in Figure 5a of an eccentric core ([CA\(^+\)]\(_{CI}\) = 0.075 mM) throughout the disassembly process provide insight into the origins of this behavior. During the plateau periods, pentamers switch between pentameric states (red capsomers in Figure 5a) and that of an incomplete hexamer (blue capsomers in Figure 5a). In this latter state, it is more likely for a CA dimer to dissociate from the lattice, which has been suggested as the rate-limiting step for viral uncoating. The resultant defect is then the site of spontaneous dissociation of CA. In comparison, we find that the fourth core ([CA\(^+\)]\(_{CI}\) = 0.500 mM) remains largely intact, which is evident from the flat profile in Figure 5a. However, fluctuations that correspond to the aforementioned structural instability of pentamers in pentamer-dense regions eventually result in core opening. Hence, all of the studied eccentric cores exhibit a tendency toward spontaneous CA disassembly due to inherent instabilities in high curvature regions with large pentamer density, which can contribute a secondary mechanism for viral inhibition.

The extent of disassembly within our simulated time-scales is considerably variable, but does suggest that increasing [CA\(^+\)]\(_{CI}\) is negatively correlated with a destabilization effect on fully enclosed mature cores. This is in contrast to some capsid stability assays and fluorescence experiments that suggest that purified cores are destabilized when treated with PF74. However, a direct comparison should not be made as canonical cores that are treated with PF74 and those that are assembled in the presence of PF74 may also be morphologically distinct, as suggested by our results above (see Figure 1). Our disassembly experiments instead seem to corroborate a capsid integrity assay that suggests that cores formed with high concentrations of PF74 have similar integrity to that of the WT. Hence, we expect inhibitory effects based on disassembly to only be relevant at lower concentrations of CIs. We further stress that the observed relationship between [CA\(^+\)]\(_{CI}\) and core stabilization is not necessarily indicative of the same relationship between [CA\(^+\)] and infectivity (as we expect the opposite to be true). Recall that we only simulate the disassembly of cores that are fully enclosed within our simulated time, which represents a subset of the cores observed in our simulations. We speculate that the fraction of malformed and eccentric cores that fail enclosure is positively correlated with [CA\(^+\)]\(_{CP}\) which would, for instance, corroborate the
increasing inhibition of infection at increasing PF74 concentration.\textsuperscript{33,26}

It is important to mention that our CG simulations have not explicitly considered the influence of host cellular factors, which impact downstream events such as trafficking, nuclear import, and integration as discussed previously. For instance, all-atom MD simulations have demonstrated that the binding of PF74 induces greater allosteric coupling between the cypA binding loop and a CA hinge region, although the implications for cypA activity remain unclear.\textsuperscript{63} Intriguingly, PF74 is able to inhibit infection even in the absence of some host cellular factors, including CPSF6 and cypA, although their presence can induce mostly cooperative (but sometimes antagonistic) influences on PF74 efficacy in a dose-dependent manner.\textsuperscript{65} However, the presence of NUP153 appears to be critical for PF74 efficacy at low concentrations.\textsuperscript{66} As NUP153 binds to the same pocket as PF74, it is conceivable that the latter competitively inhibits pre-integration complexes (PICs) from engaging with NUP153 for nuclear entry. The plateaus from our disassembly profiles in Figure S, such as when [CA\textsuperscript{+}]\textsubscript{0} = 0.100 mM, suggest that partially disassembled cores may exist as metastable states. Indeed, PF74-treated cores were recently found to undergo partial uncoating.\textsuperscript{66} We find that all CI-bound TODs remained bound to the partially disassembled capsids, as depicted in Figure Sb. Thus, in the event that viral uncoating proceeds uninhibited, a third mechanism of action is possible: the remaining CA assembly, which is likely to be part of the PIC, may consist of an elevated coverage of CI-bound CA that inhibit NUPs. A careful examination of this third mechanism of action is outside the scope of the current study, but would be a valuable subject for investigation.

On a final note, our simulations demonstrate that a CG model can aid interpretation of the anomalously broad activity of CIs observed in experiments. In the case of PF74, the binding of CIs to pre-established binding pockets stabilizes CA multimers in tubular structures and prevents disassembly.\textsuperscript{26,30,31} Our simulations incorporate this stabilization effect with the use of TODs that are fixed during dynamics. We only consider a low stoichiometric ratio of CI-bound CA given that experimental EC\textsubscript{50} estimates tend to be in the range of \(10^{-6}\) M concentrations or below.\textsuperscript{23,25} Our results show that perturbing TOD distributions in this way is sufficient to induce alternative assembly pathways that yield diverse capsid morphologies, including eccentric cores that were previously seen as a minority population in WT viruses.\textsuperscript{43–45} Given the importance of CIs during the assembly process, we suggest that future studies take care in distinguishing inhibitor behavior based on post-treatment of viral cores and integration during late stages (i.e., virion production) of the viral lifecycle. The increased diversity of capsids formed under the influence of CIs compared to WT virus results in multiple mechanisms of action, which may help explain the cooperativity that is suggested by PF74 dose–response curves.\textsuperscript{65} Furthermore, these mechanisms appear to be fundamentally different from that of inhibitors that target monomeric CA, which tend to affect either early or late (but not both) stages of the viral lifecycle. For example, CAP-1 and CAI bind to CA monomers and prevent proper CA association at key interfaces (e.g., the CTD dimeric interface) during late-stage assembly.\textsuperscript{77,80} For these types of CIs, higher drug concentrations (e.g., \(10^{-6}\) M or above) may have been required to ensure that the population of apo CA monomers is sufficiently repressed to prevent complete assembly. Hence, CIs that target the highly regulated dynamical processes observed throughout the viral lifecycle by means of CA oligomer subpopulations may have two primary benefits: (1) broader spectrum inhibitory effects and (2) reduced stoichiometric requirements that facilitate drug delivery challenges.

\section*{CONCLUSIONS}

In summary, we use CG molecular dynamics simulations and elucidate underlying mechanisms that contribute to the broad-spectrum HIV-1 inhibitory effects of capsid inhibitors (CIs), such as PF74 and GS-CA1, by virtue of small perturbations to the hierarchical self-assembly of viral capsid (CA) proteins. We find that an important mode of action by CIs is to stimulate CA association and propagate anisotropic assembly pathways, which results from the stabilization of a subpopulation of trimer of dimers (TODs); this behavior is consistent in all of our CG simulations, even at the lowest accessible drug to CA stoichiometry of less than 1 mol%. Consequently, pentameric defects appear to be rapidly incorporated, thereby increasing the expression of eccentric cores, that is, mature cores with inherently large curvature and pleomorphism. Capsids of this phenotype may unsuccessfully enclose the viral nucleic acid and nucleocapsid protein (RNP) complex, which is one possible failure mode for infection. However, in the event that the RNP complex is encapsulated, a secondary failure mode during post-entry events is observed from our capsid disassembly simulations. Here, regions of the capsid that are dense with pentameric defects appear to be inherently unstable and initial CA dissociation; eccentric cores that form due to CIs may therefore spontaneously disassemble after entry into the cytoplasm. Taken together, our findings reveal that CIs affect processes during both viral maturation and uncoating. While not explicitly observed in this work, we expect additional inhibitory effects to emerge during processes involving host cellular factors. Our results also suggest that the identification of potential binding pockets that appear upon CA oligomerization, especially those with small degrees of genetic polymorphism, and the design of small molecules that specifically target these sites—even at low stoichiometric loadings—will facilitate the design of HIV-1 (and other viral) inhibitors with similarly broad mechanisms of action.

\section*{METHODS}

Coarse-Grained Modeling and Simulation. As mentioned above, the CG models used in this work, e.g., for CA and the inert crowding agent, are described in ref 11 and summarized in Figure S1. Our simulations contained CA, pre-assembled TODs, and inert crowding agent, as described in ref 11 and summarized in Figure S1. Our simulations contained CA, pre-assembled TODs, and inert crowding agent, as described in ref 11 and summarized in Figure S1. The key difference from the simulations performed in ref 11 is the addition of pre-assembled TODs using an ENM between adjacent dimer domains (depicted in Figure S1) to preserve the inter-dimer distances around the binding pocket for CIs such as PF74 and GS-CA1; a harmonic bending stiffness of 1.0 kcal/mol was used. Configurations were allowed to equilibrate over \(5 \times 10^{6}\) \(\tau\) without any attractive interactions. Production runs were then performed as described in the main text, while [CA\textsuperscript{+}]/[CA\textsuperscript{−}] UCG state switching was attempted every \(5 \times 10^{6}\) \(\tau\), i.e., the subset of assembly-competent CA dimers from all available CA dimers in solution were stochastically chosen while a constant fraction of active species was maintained. All simulations were performed in the constant NVT ensemble at 300 K using a Langevin thermostat\textsuperscript{76} with a damping period of 100 ps and \(r = 10\) fs. All simulations were run using an in-house MD engine called UCG-MD, which is optimized to run implicit-solvent UCG simulations with dynamic load-balancing and runtime algorithms.\textsuperscript{70}
Neighbor lists up to 4 nm with an additional 1 nm skin depth were used, while load balancing was attempted every 2 × 10^6 τ. Molecular snapshots were saved every 1 × 10^5 τ for analysis.

**Data Analysis.** Graph analysis was performed using the python package NetworkX 2.1 (http://networkx.github.io/). A graph was constructed for each molecular snapshot following the criteria described in the main text and Figure S3. Hexamers and pentamers were identified by vertices (i.e., CA monomers) that participated in a closed cycle of length six and five, respectively. The remaining nodes were identified as components of incomplete capsomers. The λ of each vertex was computed using the eccentricity function.

**ASSOCIATED CONTENT**

* Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.9b01413.

Additional details on the computational models and analysis metrics used throughout this work, including Figures S1–S4 and Table S1 (PDF)

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**Notes**
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