Self-assembly of polyhedral shells: A molecular dynamics study

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The use of reduced models for investigating the self-assembly dynamics underlying protein shell formation in spherical viruses is described. The spontaneous self-assembly of these polyhedral, supramolecular structures, in which icosahedral symmetry is a conspicuous feature, is a phenomenon whose dynamics remain unexplored; studying the growth process by means of computer simulation provides access to the mechanisms underlying assembly. In order to capture the more universal aspects of self-assembly, namely the manner in which component shapes influence structure and assembly pathway, in this exploratory study low-resolution approximations are used to represent the basic protein building blocks. Alternative approaches involving both irreversible and reversible assembly are discussed, models based on both schemes are introduced, and examples of the resulting behavior described.

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I. INTRODUCTION

Self-assembly at the molecular scale is a ubiquitous process in the natural world and predicted to play a significant role in advancing nanotechnology. The formation of the protein shells, known as capsids, that provide the packaging for spherical (or polyhedral) viruses is a particularly familiar instance of natural self-assembly; the rational design of antiviral agents would benefit from an improved understanding of how such shells assemble, a phenomenon at the border between biology and chemistry. Direct visualization of evolving molecular assemblies is inherently difficult, since intermediate states elude experimental capture, and final states reveal little about their assembly pathways; thus the simulation of suitably designed models ought to prove helpful in exploring the processes and pathways involved.

Virus capsids have highly symmetric shapes, reflecting the fact that they are assembled from multiple copies of either a single molecular component—the capsomer—or a small number of distinct capsomers. Capsid assembly, a process whose details are little understood, is governed by different classes of interactions: There are interactions between capsomer proteins and nucleic acids (in the form of DNA or RNA) that initiate and regulate the assembly process, and subsequently stabilize the packaged genetic material inside the capsid. More significantly, from the perspective of the present study, there are also interactions between the capsomer proteins that stabilize the shell structure itself. What makes capsid self-assembly an ideal candidate for simulation, despite this apparent biochemical complexity, is the fact that it is able to occur reversibly in vitro (with scaffold proteins participating in the growth but not affecting stability), without the genetic material that is essential to the virus in vivo (in other cases the nucleic acid does play a stabilizing role). In addition, structurally intact empty shells occur in vitro after removal of their contents, and viruses themselves form empty capsids after removal of their background information of this kind simplifies the model design considerably, since the system need only consist of a very small number of well-characterized component types. The goal of the present work is to use molecular dynamics (MD) simulation in modeling the capsid assembly process, based on suitably simplified representations of the essential molecular components.

Motivation for simplified descriptions, that avoid becoming embroiled in the detailed physical and chemical characteristics of the capsomers, stems from the prominence of icosahedral symmetry among spherical viruses, irrespective of their biological origins. Nature has adopted this structural motif (the other basic design is a helical tube) precisely because the high degree of symmetry leads to a minimal set of construction rules. Since the task of the genetic information carried by the viral nucleic acid is not only to instruct the virus how to infect the host, but also to specify how it must replicate itself, if less information can be devoted the latter mission, more will be available for the pernicious primary task. At the same time, shapes based on icosahedra come close to maximizing the volume to surface ratio, another advantageous feature. Beyond their basic packaging role, capsomers also have an important function in the virus life cycle that the simple structural models do not attempt to address.

Self-assembly is related to crystallization, in the sense that both are governed by the laws of thermodynamics, with the obvious difference that while crystals can, in principle, grow without limit, viral growth is self-limiting. The processes are driven by bond formation between assembling units, whether atoms or protein complexes, with the goal of reaching the relevant minimal free-energy state. Assembly of symmetric structures occurs both in biological and nonbiological contexts; while crystal growth may not be of particular biological relevance, the formation of icosahedral fullerene molecules is...
an indication that some geometries are common to both. Analogous icosahedral motifs are to be found in geodesic domes; these owe their detailed structure both to the same minimalist construction rules, as well as to considerations of optimal rigidity. Nature embodies a great deal of what is considered successful engineering design, and indeed, many noteworthy engineering achievements borrow from the geometric forms found in nature, even though size scales and materials can differ greatly.

In order to avoid a substantial, if not overwhelming, degree of incorrect assembly, it seems plausible that the construction of viral capsids demands a generalized scheme independent of many of the molecular details of individual viruses; nonspecific assembly pathways might well exist, a knowledge of which could be important in finding ways to modify or inhibit the construction process, with obvious therapeutic and technological benefits. Consequently, initial exploration should focus on the basic ‘shape’ of the constituent capsomers, clearly an important factor in determining the assembly outcome. Such low-resolution descriptions amount to little more than caricatures, but, if general organizational principles do exist, they could well be sufficiently robust to reveal themselves in models from which extraneous detail has been removed. There is a longstanding tradition of working with reduced models that has proved extremely successful with other inherently complex phenomena, such as micelle growth and protein folding.

The use of MD for modeling capsid self-assembly was introduced in a previous study involving polyhedra built from 60 triangular units. In the present paper the work is extended in two directions, first by showing how more relevant trapezoidal capsomer shapes can be used and larger shells constructed, and second by demonstrating alternative ways of designing capsomer models. Despite the simplifications, such simulations provide access to assembly pathways, and are able to predict time-dependent partial structure populations that can, in principle, be compared with experiment. The following sections describe alternative model designs (Section II), the nature of the bond interactions (Section III), different assembly scenarios (Section IV), including the pathways and supplementary interaction rules involved, computational techniques (Section V), and results (Section VI) from extensive simulations. It should be stressed that while the focus is primarily on the formation of capsid shells, the method is not in any way restricted to virus structures, and is applicable to supramolecular assembly in general; a brief demonstration (Section VII) of this capability precedes the concluding section.

II. CAPSOMER DESIGN

A. General considerations

Individual capsomers are large protein complexes; owing to their size and complexity, such structures must be represented in a highly reduced form, while retaining sufficient detail to ensure meaningful behavior when studied by MD simulation. Viewed from this perspective, capsomers have two principal but not entirely independent characteristics. One is the effective molecular shape; this must be tailored to ensure capsomers fit together to form closed polyhedral shells. The other concerns the interactions between adjacent capsomer regions in the final structure; these are responsible for driving self-assembly and maintaining the structural integrity of the finished shell, and must be defined accordingly.

There are few guidelines to aid in the model design. While general thermodynamic considerations can help choose ranges for the force parameters, the approach itself is entirely empirical. Progress in this kind of discrete-particle modeling is tied to the advance in available computer power, with increasing computational capability allowing the incorporation of additional features that aid the assembly process. Other, entirely different techniques have also been employed. One considered the dynamics of spherically symmetric particles subject to directional interactions, whose states and binding energies are selected probabilistically based on rules involving local neighborhoods. Another studied disk packings on a spherical surface using a mean-field statistical mechanical analysis dependent on curvature and coverage.

No solvent is included in the simulation, an approximation borrowed from protein folding; even a neutral solvent would increase the computational effort substantially without influencing the outcome, not only because of the additional particles involved, but also because of the slower capsomer dynamics when moving in a solvent rather than in a vacuum. (In future detailed work, accounting for the effect of, e.g., pH and salt concentration on the pathways and the final state, will require more substantial models that address interfacial interactions and solvation effects at a molecular level. An alternative, Langevin-like representation of the solvent contribution as small random impulses, suitably distributed over space and time, has also not been used; if the dynamics are dominated by intermolecular forces, as is the case here, there will be little overall effect on the eventual outcome.)

The need to assure defect-free assembly allows a design tradeoff between model capsomer size and interaction complexity. Bonding of real capsomers involves the participation of a relatively large number of interaction-site pairs across a mutual contact surface. Although such interactions are not individually directional, in view of the substantial size of the capsomer compared to the effective interaction range, it is unlikely that the interactions are capable of producing strongly bound states in which capsomers are incorrectly aligned. This is not necessarily true for simplified models with few interaction sites and an overall capsomer size similar to the range of attraction, where small clusters can become trapped in states corresponding to spurious local energy minima.
instead of developing into complete shells.

One method of avoiding this situation was introduced for small capsomer models and employs rules governing the interactions (Section III) in addition to permanent bonds \( \delta \). The alternative is to use an enlarged capsomer with more interaction sites and reversible bonds; the computational cost is increased, but the enhanced structural rigidity reduces the opportunity for incorrect bonding. These design alternatives demonstrate possible approaches to the problem. They also reflect a changing perspective brought about by the fact that the available computer resources continued to increase throughout the study; greater computing power allows better capsomer representations, larger systems and longer runs, all contributing to the eventual shift from permanent to reversible bonding.

### B. Capsomer structure

The simplified trapezoidal capsomer representation employs a rigid assembly of several soft (more precisely, almost hard) spherical particles arranged in a partly overlapping configuration approximating the desired shape; specific designs are described in Section III and illustrated in Figures 11, 13 below. The repulsive force between spheres used to prevent significant spatial overlap is based on the truncated Lennard-Jones potential

\[
u(r) = \begin{cases} 
4\epsilon[(\sigma/r)^{12} - (\sigma/r)^6 + 1/4] & r < r_c \\
0 & r \geq r_c
\end{cases}
\]

where \( r \) is the sphere separation, \( \sigma \) approximates the sphere diameter, \( \epsilon \) determines the energy scale, and \( r_c = 2^{1/6}\sigma \) is the interaction cutoff; in the reduced units used subsequently, \( \sigma = \epsilon = 1 \). A suitably arranged set of spheres provides an adequate approximation to the desired shape, but as more spheres are used the computational effort required to evaluate the interactions between nearby molecules grows. The description based on pair interactions alone is, however, much simpler than alternative shape representations that require evaluating the overlap of complex rigid bodies.

Capsid size limits the amount of genetic material that can be stored. Capsids consisting of 60 capsomer units exist, but their internal volumes are insufficient for substantial amounts of nucleic acid; shells are generally larger, consisting of multiples of 60, i.e., 60T, units, where the triangulation number T can have values from 1 to beyond 25. All sizes share the icosahedral symmetry, but since 60 is the maximum size under conditions of complete equivalence, the concept of quasi-equivalence is invoked to explain how larger structures can be constructed \( \delta, 14, 13 \). Inspiration for this generalization came from the geometrical principles developed by Buckminster Fuller for geodesic domes, and construction entails arranging 12 pentamers and certain specific numbers of hexamers in a symmetric manner to produce a close approximation of a sphere that retains the 60-fold symmetry. If identical capsomer proteins are used, but a small amount of conformational deformation permitted (remaining within acceptable variations for bond lengths and angles) to achieve a minimum free energy structure, the same design principles lead to a general way of constructing shells, with the ubiquitous icosahedral symmetry as a necessary consequence.

Viewed from a simplified geometric perspective, hexamers are planar oligomers constructed out of six triangular triplets, each consisting of three trapezoidal capsomers. A nonplanar pentamer can be formed by removing one triangle from the hexamer and closing the gap. In the assembly of a polyhedral shell, the natural tendency to grow hexagons at certain positions is overcome by the more global free-energy benefits of forming a pentagon. Molecular ‘switches’, a conformation-modifying mechanism known as autostery, regulate formation of hexamers or pentamers, while ensuring these different subassemblies are positioned appropriately in the surface lattice. Without this conformational polymorphism, assembly would produce either flat hexamer sheets or T=1 polyhedra formed entirely of pentamers. In the case of quasi-equivalence, autostery represents an important characteristic of the process as it allows otherwise identical capsomers to occupy spatially nonequivalent locations in the shell. (Quasi-equivalence is only a general design consideration, and while appropriate for some viruses, may offer only a partial explanation for others; a full understanding of any given capsid structure requires analysis of the bonding energies of the capsomers themselves.)

Implicit in the model design is the fixed shape. This issue is avoided when studying T=3 assembly by defining three slightly different capsomer shapes (Section II C) destined to occupy different classes of locations within the shell. Explicit inclusion of an autosteric mechanism to provide the conformational changes required by quasi-equivalence \( 16 \) – mechanical analogs of such processes are discussed in \( 17 \) – would complicate the models.

The capsomer bonding forces are associated with interaction sites suitably positioned within the simplified structures, as shown in Figures 11, 13 below; specific pairs of sites interact whenever they approach to within a given range, and the closed-shell configuration corresponds to the minimum-energy state in which adjoining capsomers are correctly oriented; bond interactions are discussed in Section III. In a properly implemented design the only structures that should be capable of self-assembly are partial or complete shells, with energetic considerations excluding an enormous variety of ‘mutant’ structures.

### C. Specific designs

The smaller of the two cases considered here is a capsid shell of 60 identical trapezoidal capsomers. The shell can be regarded as an icosahedron \( 15 \) each of whose
20 equilateral triangular faces is subdivided into three coplanar trapezoidal units representing the capsomers of a T=1 virus, as shown in Figure 4 below. (Earlier work dealt with the simpler task of assembling 60-faced pentakisicosahedra from triangular units.) The lateral capsomer faces within the triangle are normal to the triangular plane, whereas those along the outside of the triangle are inclined at $20.905^\circ$ to the normal, resulting in a dihedral angle of $138.190^\circ$. Successful assembly is conditional upon correct relative dimensions to ensure components fit together, and angles consistent with the overall shell curvature.

The larger case corresponds to a T=3 virus. The capsid is based on a rhombic triacontahedron with 30 identical rhombic faces; each face is subdivided into two isosceles triangles (the base angles are $58.283^\circ$, so the triangles are almost equilateral), and each of these triangles is then divided into three coplanar trapezoidal capsomers yielding a total of 180; the assembled shell appears in Figure 5 below. The lateral faces within the same triangle, and between the triangles comprising the rhombus, are normal to the triangular plane, while the other faces are inclined at $18^\circ$, producing a dihedral angle of $144^\circ$. As discussed in Section II B, the use of three capsomer variants with slightly different face angles (and attractive interactions only between corresponding face pairs) avoids the quasi-equivalence issue.

Figure 1 shows the trapezoidal capsomer used in the T=1 permanent bond studies. The larger, slightly overlapping spheres that provide the overall shape occupy a single plane, while the interaction sites, represented by small spheres for visual convenience, determine the locations and orientations of the lateral faces. Each bond requires the coupling of three complementary pairs of interaction sites. The edges have between one and three interior spheres, and the triangular sets of bonding sites extend well beyond the volume of the capsomer spheres (pointing in the direction of the shell interior when the capsomer is correctly positioned). Two representations are shown; one in terms of the spheres and interaction sites, the other a block approximating the overall shape; this simplified model should be contrasted with real capsomers consisting of long, folded proteins whose exposed surfaces form relatively complex landscapes.

The corresponding T=3 version is shown in Figure 2. Since the three capsomer types differ only slightly, just one is shown, in different orientations. It has a smaller planar area than for T=1 in order to allow smaller shells relative to the size of the simulation region, but consists of two touching layers of spheres providing greater depth that helps avoid unwanted interactions. The spheres overlap within each layer, and this is varied to adjust the lateral face slopes.

The final capsomer form, appearing in Figure 3, is used for the reversible bond case with T=1. The shape is produced using three layers of spheres to reduce even further the likelihood of incorrect bonding; this is now a more important issue since there are no rules (see Section IV B) to help avoid interactions that do not contribute to the final shell. Sphere spacing within layers varies from overlapping to well-separated. Each bond now involves four pairs of interaction sites (more closely spaced than before); the energetic gain of a correctly aligned state is enhanced by distributing the interactions over more site pairs. Because of the increased capsomer thickness (three sphere layers rather than one or two) interaction sites can be positioned without extending beyond the actual area of the lateral faces, and the resulting steric screening helps prevent unwanted interactions.

Figures 4 and 5 show complete shells that these capsomers are capable of forming. (The actual size of these shells can be deduced from the ratio of icosahedron edge length to radius, namely $\sqrt{\frac{5}{3}} - 1 = 1.236 \ldots$) The former shows a T=1 shell; to make the bond locations visible the
FIG. 3: (Color online) Capsomer model used for reversible bonding (T=1); different views of the sphere-based structure and its effective shape are shown.

FIG. 4: (Color online) Complete T=1 shell (as produced by the simulation) with 60 capsomers; capsomers are shown reduced in size so that bonds are visible.

capsomers have been reduced slightly in size. The latter shows a T=3 shell formed from 180 trapezoidal units of the three different types; here capsomers are drawn to show their effective sizes hiding the bonds. Due to the nature of the bonding forces – see Eq. (2) – mutually attracting interaction sites are spatially coincident in the ground state.

The designs leave ample scope for enhancement. More complex capsomer surfaces, for example, would allow the introduction of a ‘lock-and-key’ mechanism due to steric effects. An attempt was made to use such a technique in early work on triangular units, by adding an extra sphere to the center of a lateral face and leaving an opening in the complementary face, but this mechanism did not provide the desired additional rigidity; it was not tested with larger units however, since the increased size and multiple interaction sites accomplish the same goal of restricting internal degrees of freedom.

FIG. 5: (Color online) Complete T=3 shell shell with 180 capsomers of three (color-coded) types.

III. BOND INTERACTIONS

Two fully-bonded capsomers are held together by interactions between sets of either three or four pairs of complementary interaction sites that can only interact with one another. The use of multiple sites, in addition to being more realistic, helps accomplish several goals: (a) The orientation of a lateral face and, consequently, the dihedral angle between capsomers, is specified by the plane containing the sites. (b) The overall binding interaction is distributed over the contact face; this ensures that the total binding energy of misaligned capsomers can only be a fraction of the ground-state value, thereby lessening the stability of the bond. (c) Finally, multiple binding sites enhance structural rigidity by suppressing internal modes such as twisting or flapping. Typically (though not always), two capsomers are drawn together initially by just one of the interaction site pairs, and they then reorient so the remaining site pairs can participate.

As shown in Figures 1-3, each of the three short capsomer faces contains a single set of interaction sites, while the long face contains two sets. The labeling scheme used for the sets appears in Figure 6. Since color coding is use-
ful for $T=3$, capsomers are labeled $B$, $G$, $R$ (blue, green, red) following the convention used in [4]. For $T=1$ the trimer forms an equilateral triangle (as shown), whereas for $T=3$ a small change of apex angle makes the triangle isosceles (Section II C). The five interaction sites follow the same bonding pattern in each capsomer, namely sites 2 and 3 can bond, as can 4 and 5, while sites labeled 1 bond with each other. Also shown are the site pairings associated with trimer, dimer, and pentamer or hexamer formation.

Three capsomers joined by 2-3 bonds produce a planar triangular trimer. A single 1-1 bond forms a dimer; it is nonplanar for $T=1$, while for $T=3$ it is planar when two type $G$ capsomers are involved and nonplanar in the alternative R-B case. The 4-5 bonds produce a nonplanar, flower-like pentamer for $T=1$, while for $T=3$, if all capsomers are of type $B$ they produce a pentamer, or a hexamer if alternating R and G types are involved (in three coplanar pairs).

The functional form of the attractive potential between interaction sites is a negative power of the site separation $r$, provided the sites are not too close, but for $r < r_h$ it takes the form of a narrow harmonic well; this form is chosen for convenience, and when the system is at rest $r = 0$. In the case of reversible bonding, the force is derived from the potential

$$u(r) = \begin{cases} 
  \epsilon(1/r_a^2 + r^2/r_h^4 - 2/r_h^2) & r < r_h \\
  \epsilon(1/r_a^2 - 1/r^2) & r_h \leq r < r_a 
\end{cases}$$

with typical parameter values $\epsilon = 0.1$, $r_h = 0.3$, and cutoff $r_a = 2$. The interactions used with permanent bonding are similar, although details differ slightly; earlier work also included an explicit torsional interaction to accelerate the bonding process, but this was discarded once it became apparent that pair interactions alone were sufficient.

At this juncture the two techniques diverge. In the approach used initially, bond formation is regarded as irreversible, while the alternative is to treat bonds simply as potential wells of finite depth; these represent, respectively, the extremes of kinetically limited and equilibrium assembly. Physical justification for permanent bonds stems from possible conformational changes, and even cleavage, experienced by proteins in the course of bonding. Formation of a permanent bond between two interaction sites can be implemented using Eq. (2): when the site separation initially falls below $r_h$, the potential is replaced by the harmonic term alone, irrespective of $r$, producing an infinitely high barrier from which escape is impossible. Implementation requires monitoring the identities of interacting site pairs, but this enables another feature, namely, that once a pair of sites have bonded permanently they then attract only each other, thereby reducing a tendency to form amorphous globules.

The rapid assembly associated with permanent bonding implies a more promising approach than bonds subject to breakage, but exploration reveals certain less desirable features. Stretchable bonds mean that partially formed assemblies are subject to structural distortions sufficiently large for bonds to appear between capsomers that, in more rigid assemblies, would be beyond bonding range. This leads to defects that not only prevent full assembly but also spawn mutant structures. Figure 7 shows a rare defective assembly in which growth of a sec-
ond outer layer begins at a misaligned shell element.

This situation is avoidable by making bonds nonpermanent, allowing energetic considerations to inhibit long-term survival of incorrect bond pairings. The severity of the problem can also be reduced for permanent bonds, but this requires the introduction of additional features—assembly or interaction ‘rules’—some physically motivated, others arbitrary; since similar issues are likely to appear in other kinds of self-assembly simulations, a brief discussion of the rule-based approach appears in Section IV B.

IV. ASSEMBLY SCENARIOS

A. Pathways

Polyhedral shells can assemble in many ways, even after allowing for the icosahedral symmetry. The presence of systematic features in the construction process is an important issue, since the likelihood of two arbitrary subassemblies being able to mesh successfully is low (unless incompatible pieces can be discarded in the process), but whether preferred assembly pathways exist is unknown. One hypothetical assembly scenario is based on a multistage process, with small clusters of specified shape forming initially, and then combining into increasingly larger subassemblies. These small clusters must be able to ‘tile’ the full shell, so a possible first stage is the assembly of triangles from the trapezoidal capsomers, while in the second stage these trimers are added (one at a time) to build full shells (combining subassemblies each consisting of several trimers is again an event with a low success rate). Experimental signatures of multistage assembly would be certain preferred intermediate cluster sizes, or the more accessible rate concentration dependence. In simulations that involve permanent bonding, assembly order can be enforced by using bonding rules, as described in Section IV B.

In the case of reversible bonding, where bond breakage occurs as a consequence of thermal fluctuations, assembly order can be biased by energetic preferences; in order to encourage early dimer or trimer formation, larger force constants would be associated with the relevant bonds (here, the strength is doubled). Left unattended, the system evolves to a state with many partial shells, from which further growth is impossible; to ensure there is always an adequate supply of free capsomers, partial shells below a certain threshold size are broken up at regular intervals (how often depends on growth rate) by switching off their bonding interactions for a short period. (A possible modification useful for quantitative work entails breaking up newly completed shells as well, to increase the number of shell-growth histories available for study.)

B. Interaction rules

Permanent bonds do not allow construction errors to be rectified, so it is crucial to reduce any opportunity for incorrect bond formation; this is accomplished by introducing rules governing when attractive forces may act. Analogous rules could appear in real molecules through local conformational variation in response to changes in bonding state, although their existence is not readily established; here, rules are introduced to compensate for other design simplifications. As a historical aside, the simulations started with small polyhedra, ranging from tetrahedra to 32-faced ‘soccer balls’, with interaction rules playing a prominent role; rule complexity increased for larger polyhedra with dubious chemical justification, so the rule-based approach with permanent bonding was eventually supplanted by larger capsomers that allow greater freedom in positioning interaction sites and reversible bonding to reduce construction errors.

The most basic of the rules aims to avoid bond formation in ways inconsistent with the final structure: Over the time interval starting when one of a capsomer’s interaction sites bonds with the complementary site on another, and ending when all other sites in the set are joined, these capsomers cannot form other bonds. If construction follows a pathway in which, for example, trimers initially form, which then bond into larger structures, an analogous rule applies to entire trimers. In order to minimize any adverse effects (but without attempting to mimic nature which deals with much larger systems and is not necessarily reliant on high yields), if bonding fails to complete within a prescribed interval the existing partial bond is broken; to prevent immediate rebonding newly separated capsomers must wait for a certain time before they can begin bonding again. Such rules help ensure the release of units that cannot bond completely, as in the case of two capsomers attempting to bond simultaneously along different edges of a shell opening big enough for only one.

Other interaction rules enforce assembly pathways (Section IV A), thus if growth occurs via trimer intermediates, capsomers are first required to form trimers, and only when all internal bonds are complete can these clusters associate into larger structures. Additionally, by restricting the number of larger subassemblies that can nucleate by the joining of (e.g.) two trimers—equivalent to a rate-limiting process—it is possible to ensure a significant yield of complete shells rather than numerous fragments. Typically, this limit would be set so that 50–75% of the trimers are used by the full shells, while the rest supply an adequate background concentration of construction material; a similar restriction is applied to the formation of the trimers.

Because of the intrinsic bond flexibility, the need for further rules only becomes apparent as incorrect structures are encountered; two examples will be mentioned here. When the pathway involves dimers, the appearance of unwanted bonds is reduced by requiring that after a
1-1 bond creates a dimer, the next must be a 2-3 bond between one of the dimer members and a capsomer in a growing shell, and only then the remaining 4-5 bond; since the 2-3 bond joins sites closer to the dimer center than the 4-5 bond, this restricts partially bonded units from encountering inappropriate bonding partners. For trimer assembly, the 2-3 bonds are used to build the trimer, the 1-1 bond may then join the trimer to a shell and, last of all, the more distant (from the trimer center) 4-5 bonds are allowed to form. Such constrained bonding sequences might be attributable to conformational changes as bonding progresses.

V. SIMULATION TECHNIQUES

General MD methodology is discussed in [6]; here a brief summary of the issues relevant to capsomer simulation will suffice. Interaction calculations are carried out efficiently using neighbor lists; list construction follows the procedure used for monatomic fluids. Separate lists are used for short-range repulsive forces between the spheres giving capsomers their shape, Eq. (1), and for longer-range attractive forces between interaction sites, Eq. (2). The rotational equations of motion employ standard rigid-body methods; these, together with the translational equations, are solved using a leapfrog integrator. The simulation region is bounded by elastically reflecting hard walls, implemented using short-range repulsive forces based on Eq. (1) acting normal to the surfaces; since visualization is important, hard walls avoid potentially confusing imagery accompanying periodic boundaries. In the initial state, units are positioned on a lattice and assigned random orientations and velocities; lattice spacing determines the mean density of the capsomer ‘fluid’. Simulations based on reversible bonding – which avoid the complexity associated with interaction rules – can be run on a distributed memory (message passing) parallel computer for improved performance.

Exothermal bond formation gradually heats the system; this problem is particularly acute due to the limited number of degrees of freedom capable of absorbing the excess energy (the reason being the absence of solvent and the rigid capsomer structure). Applying a weak damping force $-\gamma (\vec{v} \cdot \vec{r}) \vec{r} / r^2$ along each bond resolves this issue, where $\vec{v}$ is the relative velocity of the interaction sites and the damping coefficient $\gamma = 0.1$. Use of constant-temperature MD ensures that the overall temperature does not change despite bond formation and damping; the net effect is to transfer energy associated with internal vibration to the motion of entire clusters.

The interaction parameters are chosen for efficient self-assembly while maintaining numerical stability; there is presently no relation to experimental association energies [10]. The number density affects the outcome and must also be established empirically: Too high a value will not provide adequate space for shells to grow without mutual interference; a density that is too low will retard growth due to capsomers lying beyond their attraction range and the lack of collisions that can, in the case of reversible bonding, help break off incorrectly bound units from partially constructed shells (the densities actually used are substantially higher than in experiment).

In the case of permanent bonding, the $T=1$ simulations employ 1000 capsomers, with 13 shells allowed to nucleate and run lengths of 2–300,000 steps; for the larger $T=3$ shells the system contains 4096 capsomers, 10 shells can nucleate and run lengths are 5–800,000 steps. Bonds are allocated 4000 steps to form, and if unsuccessful the participating sites have their attractive forces turned off over the next 1000.

The reversible bonding runs for $T=1$ follow 4096 capsomers over 10 million steps, with no limit on shell nucleation. Clusters with size $\leq 30$ are broken up every 500,000 steps by turning off their attractive interactions over the subsequent 10,000 steps. The existence of a bond (insofar as cluster measurements in Section V are concerned) depends on the separation of either a single pair of interaction sites or – the more stringent requirement – on the separations of all (either three or four) pairs associated with the bond; in either case the interaction sites must approach to within a distance 0.2 ($< r_h$) to be considered paired. Finally, in reduced MD units, capsomers have unit mass, the integration step is 0.005, the temperature unity, and the number density typically 0.004.

VI. RESULTS

A. Shell analysis

The simulation results are both quantitative and qualitative in nature; a ‘snapshot’ sequence recorded at intervals of 2000 steps over the course of each simulation run provides the data needed to recreate capsid growth for post-analysis. A run can require several days of computing on a powerful workstation, but subsequent processing requires only minimal effort. Shell properties are readily measured, allowing the mean growth statistics to be analyzed, together with the behavior of individual shells (time resolution is limited by the snapshot interval, so shortlived bonds between snapshots are missed). Animated sequences providing a condensed summary of the run, in full three-dimensional detail, are also available, although static images will have to suffice here.

Establishing shell completeness requires (a) identifying a bonded set of capsomers using cluster analysis [6] with the appropriate bonding criterion, (b) ensuring their number equals the expected shell size, and finally (c) checking that each capsomer is bound to the correct number of neighbors; if all these tests succeed then, in view of the comparative rigidity of the bonds, the cluster corresponds to a closed shell. It is somewhat arbitrary whether all site pairs, or just a single pair, must be within range; the two criteria tend to complement one another. The former, more stringent criterion produces smaller clus-
ters during early growth: this is helpful when trying to visualize the emergence of partial shells from the capsomer ‘soup’, but omits the local environment to which a growing cluster might be loosely attached. The latter produces larger clusters, including big, ill-defined structures that incorporate multiple partial shells. During later stages of assembly, both criteria lead to similar results, since as shells near completion correct capsomer positioning allows all sites to participate in bonding.

Describing the nature of incomplete shells, while straightforward when the imagery is available (a nearly complete shell is readily characterized, as is a shell with a localized defect), is not obviously quantifiable; since partially formed structures, even defect-free shell fragments, have a variety of morphologies, mechanizing their classification is nontrivial. Each such structure can be represented as a bonded network (or graph), and while it is possible to determine the topology by evaluating connectivity, and the compactness by counting missing bonds, it is not apparent how such information can be utilized. Furthermore, development is not necessarily a process whereby shells grow monotonically by accretion of individual capsomers or small subassemblies, since it is also possible for larger subassemblies to aggregate and (with reversible bonding) for groups forming partial shells to break away from larger structures.

B. Growth

The most important observation regarding the overall behavior, equally applicable to both permanent and reversible bonding scenarios, is that polyhedral shells have little difficulty growing to completion, and mutant structures are highly unlikely. Partial shells tend to have few voids in their surfaces, and shells nearing completion typically have only one or two holes; more open cagelike structures with multiple lacunae are not encountered. The results also reveal, not surprisingly, that an inappropriate choice of interactions (or, for that matter, even a slight error in defining capsomer geometry) leads to a wide variety of alternative structures, including open networks, incorrectly linked assemblies of shell fragments and amorphous shapes that defy characterization. Due to the difficulty of describing anything other than a correct shell, this aspect of the subject is avoided, but real viruses experience analogous effects in unfavorable environments.

The main emphasis of the analysis is on reversible bonding, the likely focus of future work, but a few key results for the permanent bonding alternative are presented first in order to demonstrate its capability. Figure 8 summarizes the number of complete shells as a function of time for several different cases, namely $T=1$ shells grown using monomer and trimer pathways, and $T=3$ shells using dimer and trimer pathways, with two examples of the former to show the variation between runs. Figure 9 shows several snapshots of $T=3$ growth using a trimer pathway; shells are isolated from their milieu using automated cluster analysis.

Figure 10 summarizes the yield of complete $T=1$ shells vs time for simulations employing reversible bonding, three runs each for unweighted, dimer- and trimer-weighted pathways; note that the runs are now an order of magnitude longer. Dimer growth appears to produce the highest yields at intermediate times, but any observation of this kind is tentative because the spread in trimer results over the runs suggests more sampling is required and also because the effectiveness of selectively increased interaction strengths in biasing the pathways has yet to be established. Growth snapshots using trimer weighting appear in Figure 11.

C. Shell statistics

Despite the difficulty in quantifying shell growth, average properties of partial shells can be used in studying growth trends. Capsomers belonging to incomplete shells can be categorized according to the degree of bonding; a higher number with the maximal five bonds is consistent with a more compact hole-free partial shell, whereas an increased number with just one or two bonds is an indication that at least part of the structure is loosely connected. Figures 14 and 15 show the capsomer fractions as functions of shell size for unweighted and trimer-weighted assembly; averaging is over the entire run and the stricter (all-pairs) definition of bonding is used. Two differences are apparent; in the unweighted case there is a larger proportion of singly bonded units in the small (size $< 20$) clusters, while in the trimer case there is an enhanced preference for five bonds over four in the size range 30–50 (possibly reflecting a drop in monomer-sized holes in the shell); both of these are indicators that trimer weighting leads to more compact intermediate states (dimer results are similar).

Figure 12 shows how the size distribution develops with time; prominent features are the steady trend to comple-
FIG. 9: (Color online) Snapshots from a T=3 simulation with permanent bonding: the images show an early state (including the container for reference); three views at different times showing only the growing shells; the entire system corresponding to the third of these views; the final state (capsomers are colored by type).

FIG. 10: Number of complete shells vs time for reversible bonding.

dition, a relatively narrow distribution once growth is well underway, and the imposed cluster breakup. The fact that for most of the run there are few clusters of intermediate size is a consequence of breakup followed by prompt attachment of newly freed units to larger assemblies. At the end of the run the system consists entirely of shells that are either complete, or nearly so, together with monomers and small fragments.

Detailed information extending beyond such system-wide averages is obtained by examining individual shell growth; two forms of analysis are demonstrated here. The first ranks the clusters in each snapshot configuration by size and then graphs the size history of the largest of these clusters, the 5th largest, and so on, until the count reaches the number of shells that grow to completion; no attempt is made to ensure ‘continuity’ by tracking specific clusters. The results, using the strict all-pairs bond definition, appear in Figure 15. Figure 16 shows the corresponding results when only a single pairing is required, leading to larger clusters early in the run. The long-term behavior in both cases is the same (graphs terminate when shells are complete). The periodic breakup affects small clusters until their size exceeds the threshold; there is a secondary effect on larger clusters since newly freed capsomers provide competition as bonding partners.

The second kind of analysis tracks, as closely as possible, the development of particular shells. This is accomplished by first identifying all complete shells in the system at the end of the run, together with their constituent units, and then using this information while following the construction history of each shell. It is optional whether to include capsomers whose shell membership is transient, an effect that appears early in assembly, but less so later on when a high degree of bonding makes escape more difficult. Ambiguity can arise as to a shell’s true ancestry; where there are several contributing precursor assemblies, it is the one most heavily represented in the final shell that is credited.

Figure 17 shows typical shell growth histories for a trimer-weighted pathway. The selected shells demonstrate different growth characteristics; they include both the fastest shell to complete, and one of slowest, taking
FIG. 11: (Color online) Snapshots from a simulation with reversible bonding; early state; two views after $0.5 \times 10^6$ steps showing the full system and the 132 clusters of size $\geq 10$ (color coding - red particles have all five bonds in place, blue have $< 5$); the 34 clusters of size $\geq 50$ after $2.5 \times 10^6$ steps; two views of the final state after $8 \times 10^6$ steps showing the entire system, and just the 39 complete shells with container.

FIG. 12: Capsomer fractions with different bond counts vs cluster size for reversible bonding and unweighted assembly.

FIG. 13: As in Figure 12 for trimer-weighted assembly.

some three times longer, although in most cases a delay beyond about two million steps is caused by a wait for the final few capsomers (possibly just one) to bond. The strict bond definition is used and histories end upon shell completion; the almost monotonically increasing solid graphs show how many members of the final shell have already joined, while the more variable dashed graphs include transient members that eventually break away from the shell.

Visualization is extremely useful for probing the details of the behavior; unfortunately its rich message cannot be transferred to the printed page. Suitable color coding, based on known final shell membership, allows the identification of units destined to join a particular shell, as well as transient members. Capsomers, literally, can come and go; only when a capsomer is bound to most of its neighbors, and embedded in a substantial shell fraction, is it unlikely to be knocked out of position. Population exchanges of this kind are not readily characterized in a quantitative manner.
VII. OTHER SELF-ASSEMBLY EXAMPLES

Although the focus of the paper is on polyhedral shell growth, with the goal of modeling capsid formation, the same approach works for other self-assembling systems. Two examples of this kind will be mentioned briefly, the first broadly related to micelles, the second a unique and somewhat improbable structure that solves a well-known assembly puzzle. While the former consists of large number of identical components, the latter has highly specific interactions between a small set of distinct components. The motivation underlying these examples is to show what can be accomplished in the simplest of self-assembly simulations. Related ‘analog’ simulations have been performed in the laboratory, using millimeter-size plastic objects in solution with appropriate adhesive-coated surfaces.

The first of the systems is a fluid of rigid particles (micelle simulations generally use flexible chain molecules) each having the form of a tapered cylinder made from a linear array of spheres of decreasing size. Lennard-Jones interactions occur between spheres occupying equivalent locations in different particles. The final state of an 8000-particle simulation under conditions of gradually reducing temperature (a way of encouraging collapse into the ground state) is shown in Figure 18; it consists of 28 packed spherical clusters with sizes ranging from a maximum of 368 down to 260, one hemispherical cluster of size 183 (half the maximum size), and a few monomers and tiny clusters. The range of cluster sizes is determined by how many particles of a given shape can form a packed shell, with the absence of well-characterized bonding patterns accounting for the size variability, just the opposite of what occurs with polyhedral shells.

The second system is an MD realization of the ‘Soma cube’. This is an assembly puzzle consisting of seven distinct pieces, each formed from three or four unit cubes joined along their faces in various fixed configurations; the pieces can be assembled to form a cube of edge three in 240 different ways. For the simulation, the puzzle pieces are replaced by rigid bodies made of spheres located at the cube positions, with attractive forces between spheres that are adjacent when all pieces fit together in a particular solution of the puzzle. If the bodies are randomly placed in a large container (with reflecting walls as before), assigned random initial velocities, their dynamics simulated using MD, and subjected to a gradually falling temperature, will the puzzle solution emerge spontaneously? The answer is a qualified yes; in a substantial proportion of runs the cube assembles itself. Figure 19 shows the components and the assembled product, providing yet another example of how, if interactions are formulated properly, the outcome is correct self-assembly.

VIII. CONCLUSION

The approach described in this paper involves placing simplified capsomer elements in a container and fol-
FIG. 17: Size histories of selected clusters that grow into complete shells (see Section VI C).

FIG. 18: (Color online) Tapered cylindrical particles that have condensed into spherical clusters.

Following their dynamics, the outcome is a demonstration that a simple potential energy function, based on structural considerations, is essentially all that is required to drive the assembly of the corresponding polyhedral capsid shells. The surprising aspects of the results – given the absence of any \textit{a priori} theoretical expectation – are the fast growth rates, high yields, and the avoidance of incorrect structures. Although the models are not representative of real molecules, if general principles underlying capsid assembly do exist, simplified systems of this kind ought to embody their essence.

The advantage of the simplified approach, once its validity and relevance are confirmed, is that it allows exploration of the salient features of the problem free of peripheral detail; the influence of shape and interactions can be examined relatively easily, in contrast to the heavy computational demands of an explicit atomic representation. While the qualitative benefits are indisputable, the approach is not intended for quantitative estimation,
where accurate molecular structure and interactions are likely to be important. This does not preclude basing systematic studies on simple models; it is possible, for example, to explore the effects of varying relative interaction strengths to match known binding energies, or to introduce design modifications aimed at better approximating major structural features of the capsomers. Such projects await future consideration.

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