Simulation Studies of the Self-Assembly of Cone-ShapedParticles

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We investigate the self-assembly of anisotropic cone-shaped particles decorated by ringlike attractive “patches”. In a recent paper, we demonstrated that the self-assembled clusters, which arise due to the conical particle’s anisotropic shape combined with directional attractive interactions, are precise for certain cluster sizes, resulting in a precise packing sequence of clusters of increasing sizes with decreasing cone angles (Chen et al. Proc. Natl. Acad. Sci. U.S.A. 2007, 104, 717–722). Here we explore the dependence of cluster packing on the cone angle and cooling rate and discuss the “stability” and “metastability” of the resulting structures as well as polymorphism of non-“magic-number” clusters. We investigate large clusters of cones and discuss the implications of our simulation results in the context of the Israelachvili packing rule for surfactants and a recent geometrical packing analysis on hard cones in the limit of large numbers of cones.

1. Introduction

Nature has mastered the self-assembly of building block subunits into complicated structures with extraordinary precision and accuracy. Examples range from the assembly of complementary strands of DNA into a double helix to proteins self-assembling into spherical virus capsids with icosahedral symmetry to the formation of filaments, fibers, and membranes in cells. In these examples, nature exploits the use of highly specific and directional interactions to achieve precisely ordered structures. Mimicking such interactions in synthetic nanostructures by exploiting anisotropy in both building block shape and interbuilding-block interactions may lead to the fabrication of precise structures for use in a range of applications.

This work is inspired by recent experimental interest in the assembly of conelike particles and molecules. Examples include the assembly of cone-shaped amphiphilic dendro-calixarenes into precise micelles,1 middle-functionalized conelike peptide amphiphiles forming nanofibers,2 and self-assembled superstructures by rodlike metal—polymmer amphiphiles.3 Additionally, some capsomers (morphological subunits composed of groups of proteins) in virus protein capsids4,5 possess short, truncated conelike shapes and, to a first approximation, can be modeled as cone-shaped particles. On larger scales, novel synthetic methods have been developed to make conical colloids. Sheu et al. fabricated uniform nonspherical particles from homo-IPNs (interpenetrating polymer networks), including ice-cream-cone-shaped particles via seeded emulsion polymerization of styrene—divinylbenzene mixtures in cross-linked monodisperse polystyrene seed latexes.6,7 Most recently, Doulliez8 demonstrated the synthesis of micrometer-sized hollow cones of controlled angles via self-assembly in bolaamphiphile/hexadiamine salt solutions. A recent computer simulation study showed the formation of a gyroid cubic phase by tapered or pear-shaped particles with only repulsive interactions.9 Although progress has been made in the preparation of conical building blocks, the principles underlying their self-assembly into ordered structures are not well understood. As such, it is desirable that a general, predictive theoretical or simulation approach be formulated to study this problem and provide design principles for the assembly of conical building blocks.

A second motivation for this study comes from theoretical considerations of the conical particle packing problem. In an attempt to develop guidance for the assembly of amphiphilic surfactants, Israelachvili10 described the conditions under which spherical micelles or cylindrical micelles will form, on the basis of a “critical packing parameter” (CPP) or “shape factor”. The CPP is defined as $V_{\text{dil, c}}$, where $V$ is the volume of the amphiphilic particle or molecule, $a_0$ is the surface area of the head group, and $l_c$ is the critical length. According to the Israelachvili packing rule, when $CPP \leq 1/3$, as in ideal cones, amphiphiles assemble into spherical micelles and when $1/3 < CPP \leq 1/2$, as in truncated cones, cylindrical micelles form. Though the Israelachvili packing rule is successful in rationalizing the self-assembled structures formed in surfactant systems, the rules provide primarily qualitative insight and has limited application. These limitations were recently addressed by Tsonchev et al., who developed a geometric packing analysis for the packing problem of amphiphilic nanoparticles treated as hard cones.11 With the


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assumption that all particles pack locally in a hexagonal arrangement, Tsonchev et al. predicted that spherical micelles are always preferred in the self-assembly of hard conical particles because of the higher packing fraction in spherical clusters than in cylindrical clusters, in contrast to the commonly held belief that truncated cones will form cylindrical micelles. However, the key assumption of local hexagonal packing in Tsonchev et al.’s theory is only valid in the limit of very large N, where N is the number of cones in a single cluster. For clusters comprised of smaller numbers of particles, the finite curvature of the assembled cluster will prevent extended hexagonal order, altering the cluster shape in an unknown way. For example, as we will show, twelve cone-shaped particles with an angle of 62° form a perfect icosahedral cluster where every cone base has only five nearest neighbors (pentagonal packing) instead of six (hexagonal packing). Furthermore, both the Israelachvili packing rule and the geometric packing analysis of Tsonchev et al. focus on the explanation of overall cluster shape and not the local packings of the particles within the clusters.

Recently, Rapaport13 reported a molecular dynamics (MD) study on the self-assembly of polyhedral shells to investigate the dynamics underlying protein shell formation in viral viruses. He demonstrated an extension of his method to study rigid tapered cylindrical particles (i.e., cones) and showed several large spherical micelles formed by many tapered cylindrical particles in MD simulations. The present simulation work focuses on the local packing for small to intermediate cluster sizes and also examines the validity of the above predictions for the packing of conical building blocks at a large cluster size limit.

In previous work,14 we showed that the self-assembled structures from cone-shaped particles belong to a more general packing sequence that includes polyhedral structures formed by evaporation-driven assembly of colloidal microspheres15 and several virus capsid structures. In this paper, we extend our investigation and tackle several issues unaddressed in our previous work but critical for materials design and fabrication, such as the influence of the cone angle on cluster packings, the cluster size distribution and yield, and the influence of the cooling rate on the assembled structures. Additionally, we discuss the implications of our simulation results as they pertain to the Israelachvili packing rule for surfactants and the geometrical packing analysis on hard cones.

2. Model and Simulation Method

Rigid cone-shaped particles are constructed by fusing together a linear array of overlapping spherical subunits, or beads, with decreasing sizes. Varying the number of beads and/or the interbead distance controls the length of a cone, and the cone angle is controlled by varying the size gradient along the axis of the array.

In this work, we consider six-bead cone-shaped particles in which the distance between neighboring beads within a cone is 0.5σ, where σ is the diameter of the smallest bead. As shown in Figure 1, the cone angle θ can be calculated from the equation sin(θ/2) = (r_s - r_e)/2.5, where r_s and r_e are the radii of the largest and smallest beads in the particle, respectively. The model cone can be well approximated by a truncated cone with two hemispheres attached to each of its ends. The volume can then be calculated by V = π(r_s^2 + r_e^2 + r_s r_e)h/3 + π(r_e^3 + r_s^3)/3 with h = 2.5, the distance between the centers of the two end beads. Square-well interactions of well depth ε exist between beads occupying the same positions (indicated by the same colors in Figure 1) on the particles except for the two end beads. The end beads, and unlike beads with different colors, interact through a hard-core excluded volume interaction only. This choice of interactions promotes the lateral head—head and tail—tail packing of cones, and the use of nonattractive end beads promotes the formation of closed, terminal structures and avoids cluster—cluster aggregation at dilute volume fractions. The interaction range (width of the square well) λ is fixed at 0.4σ. The particle-based simulation approach we employ here has the advantage that it makes no a priori assumptions regarding local packing.

Monte Carlo (MC) simulation in the canonical ensemble using the standard Metropolis algorithm is used to simulate the systems. The simulation box is cubic, and periodic boundary conditions are implemented in all three Cartesian coordinates. At each MC step, either a translational or a rotational move is attempted for each particle,16 and the trial move is accepted by comparing the Boltzmann factor, which is calculated from the potential energy difference between the configurations before and after the trial move, to a computer-generated random number. The ratio of attempted translational moves to rotational moves is 2:8. Initially, all cone-shaped particles are randomly distributed throughout the simulation box at a high temperature. The maximum translational displacement is chosen to be 0.2σ, and the maximum rotational displacement is chosen to be 0.2 rad. We choose a cooling rate that is affordable slow yet fast enough to produce stable structures reliably and efficiently. The influence of the cooling rate is also investigated. The system is slowly cooled to a low, target reduced temperature (0.3—0.5) to allow sufficient time for the particles in the system to assemble into ordered structures and until the monitored total system energy stops changing or the change is unnoticeable within a reasonable simulation time. All temperatures are quoted in reduced units in terms of ε/κ_b, where κ_b is the Boltzmann factor. The cone angle is varied systematically (with small increments 0.5—1°) in successive simulations to study the dependence of cluster packings on the cone angle. In addition, multiple independent runs at certain cone angles are performed with different initial configurations and along different cooling paths to investigate the path dependence of the structures. The cluster size distribution information is collected by using a previously developed analysis code.17 It should be pointed out that the stochastic nature of the Monte Carlo method, in which the particles translate and rotate diffusively, mimics the kinetics of Brownian particles but does not necessarily reproduce actual assembly kinetics. In this work, however, we are interested in equilibrium structures and not kinetics, and this motivates the use of the MC method.

![Figure 1. Illustration of the model sno-cone shaped particle.](image)


Since we are interested in the structure of clusters formed by assembly of conical building blocks, we restrict our studies to the dilute regime where individual clusters are likely to form. At high volume fractions, other structures, such as cylinders as in surfactant systems, or even a glass transition is possible, but this lies outside the scope of the present work. In the dilute regime, we find that cluster formation is determined solely by the cone angle and the particle concentration has little or no influence on cluster formation. In the simulations reported in this work, we used $M = 500–1000$ particles, and we varied the simulation box size $L$ to keep the volume fraction between 0.1 and 0.3. The volume fraction is defined as $MV/L^3$.

3. Results and Discussion

This section is organized as follows. In section 3.1, we describe the structural characteristics for each precise cluster we obtain at certain “magic number” values $N$. In section 3.2, we describe how we determine the cone angle range for each precise structure and discuss the stability of clusters on the basis of the cone angle range. Section 3.3 discusses the cluster size distribution. Section 3.4 discusses the influence of the cooling rate on the cone angle range and cluster size distribution. Finally, we discuss our simulation results for a large cluster size in the context of previous theoretical studies in section 3.5.

3.1. Cluster Structures at Small to Moderate $N$. As we gradually vary the cone angle, we find upon cooling that a series of unique clusters are obtained with a specific number of cones, $N$. Each size $N$ cluster has a range of cone angles over which that $N$-particle cluster can be assembled. Figure 2 lists the visual images of structures at $N = 4–17, 20, 27, 32,$ and $42$, along with the corresponding cone angle ranges. The collection of such information results in a diagram for the assembly of cone-shaped particles, as shown in Figure 3. The cooling rate used is $\Delta T = -0.01$ per 0.5 million MCS (Monte Carlo steps). The influence of the cooling rate on the cone angle ranges is discussed later in section 3.5. The packings observed in different simulations are robust and independent of the cone angle.

The directional attraction between cone-shaped particles is the driving force for assembly. Particles attempt to maximize their number of nearest neighbors to achieve the lowest energy state possible at the target temperature. At the same time, the conical excluded volume forces the particles to assemble into a curved, closed structure. The specific lateral attractions between cones expedite their assembly into tightly bound clusters. Once in the cluster, the outermost beads of the cones are effectively confined on the surface of a convex shell formed by the inner beads of the cones. Figure 2 shows clusters with small sizes ($N \leq 17$) have distinct polyhedral convex shapes.

Platonic solids, packings of spheres, low-energy Lennard-Jones clusters, and equilibrium configurations of point charges on a spherical surface all relate to different types of polyhedral cluster structures. While our precise packings do share some common features with some of these polyhedral structures, they also demonstrate some interesting, unique characteristics. For example, if we take the head bead (the largest bead) in a cone as a vertex and construct a convex hull formed by all vertices (head beads) within a cluster, clusters with size $N = 4–10$ and 12 shown in Figure 2 are the eight convex deltahedra with $4, 6, 8, 10, 12, 14, 16,$ and $20$ faces, i.e., tetrahedron ($N = 4$), triangular dipyramid ($N = 5$), octahedron ($N = 6$), pentagonal dipyramid ($N = 7$), snub disphenoid ($N = 8$), triaugmented triangular prism ($N = 9$), gyroelongated square dipyramid ($N = 10$), and icosahedron ($N = 12$). A deltahedron is a polyhedron whose faces consist of only equilateral triangles that are not in the same plane. The same packings were also observed in experiments on colloidal microspheres that formed clusters upon droplet evaporation; in that paper the relationship of convex deltahedra to other applications was discussed. The investigation of evaporation-driven assembly of colloidal spheres was discussed in our previous work. At the smallest possible cone angle for the $N = 8$ clusters, we observed a twisted-square structure instead of the snub disphenoid. This twisted-square structure was observed in evaporation experiments by Cho et al., and while the snub-dishpenoid structure was observed in experiments by Manoharan et al. and Yi et al.

The $N = 11$ cluster is, however, special. It is not a deltahedron since no 18-face deltahedron exists. Since one of its vertices is shared by six triangles, there should be a hexagonal packing plane if all six triangles are equilateral. This is clearly not the case for the $N = 11$ cluster as shown in Figure 2. Through the vertex that has six neighbors, there is a vertical reflection plane. Similar to the $N = 11$ cluster, the $N = 13$ cluster also has a vertical reflection plane through the cluster center. Both clusters lack high symmetry as compared to the $N = 12$ cluster with icosahedral symmetry.

The $N = 14$ cluster has a biplanar structure with $D_h$ symmetry ($D_h$ symmetry refers to dihedral rotational symmetry, or rotational symmetry with mirror symmetry) with some additional subtlety. Each plane of the $N = 14$ cluster contains seven hexagonally arranged cone-shaped particles, but with opposite rotation directions and two planes in the staggered conformation. In one plane, seven cone-shaped particles rotate clockwise with respect to the center particle into the $R$ (rectus) configuration, while in the other plane they rotate counterclockwise into the $S$ (sinister) configuration. This same packing was also observed in the evaporation-driven assembled structures of 14 colloidal microspheres but without the inherent chirality observed here that arises from the conical particle shape.

The $N = 15$ cluster has 3 hexamers, or hexavalent vertices, and 12 pentamers, or pentavalent vertices. The cluster consists of three hexagonal faces on which each hexamer is surrounded by six pentamers. The $N = 16$ cluster possesses tetrahedral symmetry and the shape of a truncated tetrahedron. The cluster has four hexamers at the center of each of its four hexagonal faces. All other 12 vertices are pentamers. The $N = 17$ cluster has 5 hexamers and 12 pentamers and possesses two vertical reflection symmetries through the top hexamer, as shown in Figure 2.

The $N = 20$ cluster has a short cylinder-like shape. It can be best described as three adjacent layers stacked together with $D_{3h}$ symmetry ($D_{3h}$ symmetry refers to dihedral rotational symmetry with reflective symmetry in a horizontal mirror). Each of the top and bottom layers contains seven particles arranged hexagonally in an eclipsed conformation, and the middle layer is a six-particle ring in a staggered conformation with both top and bottom layers.

The $N = 27$ cluster is hollow and has $D_{3h}$ symmetry. The 12 pentamers are distributed into a 1:5:5:1 arrangement and have five particles in the horizontal reflection plane, where each pentamer is surrounded by five hexamers. The two halves are

in an eclipsed conformation. This cluster structure may correspond to a nonicosahedral spherical virus structure, such as the middle component of the pea enation mosaic virus, the top component of the tobacco streak virus, and the Tulare apple mosaic virus, which consist of roughly 150 subunits or 27 capsomers (12 pentamers and 15 hexamers) and violate the Caspar-Klug (CK) quasi-equivalence theory, as suggested by Cusack. In contrast, the \( N = 8 \) cluster, which is also hollow, has icosahedral symmetry with 12 pentamers and 20 hexamers. Viruses with a triangulation number \( T = 3 \) have 32 capsomers in their capsid and exhibit the same icosahedral symmetry as shown here. \( C_{60} \) fullerenes also have the same symmetry that consists of 12 pentagonal faces and 20 hexagonal faces.

The \( N = 42 \) cluster, which is hollow and is the largest precisely packed structure that we obtain within a reasonable computation time, does not have the expected icosahedral symmetry. Interestingly, the \( D_{60} \) symmetry of this nonicosahedral structure is also predicted in the proposed optimal packings of the maximum volume of a convex hull for a set of points on a sphere—a

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closely related mathematical problem—and in the packings of point charges on a sphere, known as the Thomson problem in mathematics.\(^{24}\)

It is worth noting that all the packings at the discussed values of \(N (N = 4 - 17, 20, 27, 32, \text{ and } 42)\) in the presented sequence are unique and robust; therefore, we refer to them as magic number clusters, in analogy with Lennard-Jones clusters.\(^{25}\) There is rare or no fluctuation once the precise clusters are formed. Clusters of size \(N\) other than these values exhibit multiple polyomorphs and hence are not unique and may oscillate between different structures along a long simulation.

3.2. Cone Angle Ranges. 3.2.1. Method Used to Identify the Cone Angle Ranges. We estimate the range of angles, defined as the difference in the upper limit and lower limit of cone angles that will form clusters of a given size, by analyzing the cluster size distribution at each investigated cone angle. The upper limit of the cone angle capable of producing a cluster of a given size can be directly estimated from the cluster size distribution. Since we vary the cone angle in small increments (about 0.5–1\(^{\circ}\)), it is easy to locate the largest angle at which a given cluster size disappears. We then take the last cone angle that still yields this cluster size as the upper bound on its cone angle range.

The determination of the lower limit of the cone angle capable of producing a given cluster size is less straightforward. At these angles the system is usually a mixture of complete clusters and incomplete clusters that will eventually form slightly larger clusters. We then use visual inspection combined with the information deduced from the cluster size distribution to determine the lower limit of the cone angle that can self-assemble into clusters of a given size. Again, we take the smallest possible cone angle that still can form a given size of cluster as the lower boundary of its cone angle range.

The error in determining cone angle boundaries for a cluster of given size is limited only by the angle steps we used (0.5–1\(^{\circ}\)) in the simulations. Statistical runs for typical clusters \(N = 11–13\) demonstrate that the fluctuation of cone angle boundaries determined by this method varies within 1\(^{\circ}\), as shown in Figure 3.

The simulation-estimated range of cone angles for each structure, which can be considered as a kind of “tolerance”, shows whether it is possible for cone-shaped particles with a given angle to assemble into clusters of a given size. This information should be useful for designing materials from such cone-shaped building blocks.

3.2.2. “Stable” and “Metastable” Clusters. The range of angles over which a given magic number cluster structure is obtained is an indication of the stability of that structure at the current set of simulation parameters, and thus we can categorize the \(N\)-particle clusters as stable and metastable clusters according to their cone angle ranges. We note that this classification, while quantitative, is subjective and not based on evaluation of the cluster free energy.

Figure 3 shows that the angle ranges can differ significantly and change in a discontinuous way with respect to \(N\). The overall tendency is for the angle range to decrease as \(N\) increases. However, some clusters have unusually small or large angle ranges as compared to adjacent ones. For example, the \(N = 4\)
and 6 clusters have the widest angle ranges of 51.2° and 28.4°, respectively, while both $N = 5$ and 7 clusters have angle ranges of only about 10°. Similarly, the cone angle ranges for the $N = 9$ and 12 clusters are significantly larger than the adjacent $N = 11$ and 13 clusters.

The potential energy part of the free energy decreases monotonically as $N$ increases, as the cone particles in larger clusters have more neighboring particles of attractive interactions on average. However, entropy is not a monotonic function of $N$. The effect of entropy on the cluster stability may be understood from symmetry arguments. An icosahedron possesses 6 fivefold, 10 threefold, and 15 twofold rotational symmetries through its vertices, faces, and edges, respectively. Such high symmetry represents high configurational entropy and low free energy, assuming relatively unvaried or insignificant potential energy. As such, a 12-particle cluster with an icosahedral symmetry is particularly stable and can be found assembled from conical particles with a wide range of angles. In contrast, the $N = 11$ cluster contains one particle fewer than the most stable icosahedron cluster and, although its energy is only somewhat higher, is of substantially lower symmetry and thus lower entropy. As such, the $N = 11$ cluster has an unusually small cone angle range, indicating low stability.

In summary, we categorize the clusters into stable (such as $N = 4, 6, 9,$ and 12) and metastable (such as $N = 5, 7, 11,$ and 13) clusters, characterized by wide and narrow cone angle ranges, respectively. The configurational entropy is likely to be the cause of the discontinuous change in stability with respect to the cluster size $N$.

### 3.3. Cluster Size Distribution and Yield

The cluster size distribution gives important information about the yield or the possibility of obtaining clusters of a certain size as well as the relative stability among clusters with comparable sizes at a given cone angle. Figure 4 shows typical cluster size distributions at cone angles of 67.4° and 62°, which are optimum angles (which give the highest yield of a specific cluster size at a cooling rate of $\Delta T = -0.01$ per 0.5 million MCS) for $N = 11$ and 12, respectively.

Since the $N = 12$ cluster has extremely stable icosahedral symmetry, the size distribution at the optimum cone angle, 62°, is essentially monodisperse, as shown in Figure 4b. In contrast, even at its optimum angle the yield of the $N = 11$ cluster is still smaller than that of other clusters such as the $N = 9$ and 12 clusters at the same angle, as shown in Figure 4a. Actually the $N = 12$ icosahedral cluster is favored over the $N = 11$ cluster under all conditions. The particularly small angle range and yield of the $N = 11$ cluster indicate its low stability compared to that of other cluster sizes.

As we decrease the cone angle $\theta$, more cone-shaped particles can self-assemble into clusters with larger sizes. The difficulty of obtaining complete clusters in the simulation usually increases as $\theta$ decreases, i.e., as the cluster size increases. As such, stable clusters with small sizes, such as $N = 4$ and $N = 6$ clusters, display monodisperse distributions within a range of angles (102°–116.4° for the former and 81°–92° for the latter). In contrast, metastable clusters such as $N = 5$ and $N = 7$ clusters have a relatively broad distribution and low yield even at their optimum angles. Other stable clusters, such as the $N = 8$ and $N = 9$ clusters, also have a narrow distribution and relatively high yield (over 70%).

To conclude this part, stable clusters such as $N = 4, 6,$ and 12 clusters have a nearly monodisperse distribution and high yield in contrast to a relatively broader cluster size distribution and low yield for metastable clusters such as $N = 11$ and 13.
clusters, consistent with their corresponding wide and narrow angle ranges, respectively, in the previous discussion. In the evaporation experiments on colloidal microspheres,\textsuperscript{15,21} the cluster size is set by the number of particles trapped in a droplet. There is no direct control over the structures as they form, and separation processes are required to obtain clusters of a particular size. In contrast, with a properly chosen cone angle it is possible to control how many particles can assemble into a single cluster and with predetermined, desired precise packing. We have also predicted that precisely packed structures with a narrow size distribution can be assembled from cone-shaped particles with certain cone angles, which is of technical significance because it may substantially reduce the cost of, if not eliminate the need for, the separation procedures required in assembly of isotropic particle building blocks. This demonstrates the possibility of using patterned cone-shaped particles as programmable building blocks for self-assembly.

### 3.4. Influence of the Cooling Rate

#### 3.4.1. Influence of the Cooling Rate on the Size Distribution Profile and Cluster Yield

To investigate the influence of cooling rates, we compare the size distributions of cone-shaped particles at certain cone angles for three cooling rates. Cooling rate 0 decreases the temperature by 0.02 per 0.5 million MCS, cooling rate 1 decreases the temperature by 0.01 per 0.5 million MCS, cooling rate 2 decreases the temperature by 0.002 per 0.5 million MCS, and cooling rate 3 decreases the temperature by 0.001 per 1 million MCS.

Within certain angle ranges of stable clusters, slower cooling rates show little or no influence on the already nearly monodisperse size distribution, such as 102.5°–116.4° for the N = 4 cluster, 81°–92° for the N = 6 cluster, and 57.4°–62.7° for the N = 12 cluster.

For other cone angles, a slower cooling rate shifts the size distribution to larger N and increases the yield of larger clusters, as shown in Figure 5 for θ = 67.4°. Ideally we should use a cooling rate as slow as possible, but due to the computational limitation, we chose cooling rate 1. One relevant question is the following: Will the size distribution at all angles eventually evolve into a single sharp peak with a slow enough cooling rate? The simulations using the slowest cooling rate (cooling rate 3) still show the coexistence of at least two clusters with comparable sizes, though the slower cooling rate does appear to result in fewer peaks in the size distribution profile.

Again, we emphasize that all the packings at each magic number N are robust and are independent of the cone angles at which they are observed, e.g., 14-particle clusters at 50.3° and 55.7° are identical, although the yield of the cluster for each angle may change with different cooling rates.

#### 3.4.2. Influence of the Cooling Rate on the Cone Angle Ranges

The cone angle ranges for clusters of size N = 6, 9, 12–14, and 20 may be calculated from simulations using different cooling rates. The results are listed in Table 1.

For stable magic number clusters, we observe that the cooling rate has little or no influence on the width of their cone angle ranges. At each cone angle investigated, slow cooling shifts the size distribution to larger N and produces a higher yield for larger clusters. Since slow cooling shifts both the lower boundary and upper boundary by 0.5°–3° in the same direction, the net effect is that the cone angle range remains roughly the same, with an overall shift toward larger N.

For metastable magic number clusters, the influence of the cooling rate is, however, complicated. We find that a slow cooling rate favors those clusters that are thermodynamically more stable, such as the N = 4, 6, and 12 clusters, and suppresses, or even diminishes, the appearance of metastable clusters, such as the N = 11 and 13 clusters, motivating the use of the term metastable. For example, simulations using cooling rate 2 significantly decrease the yields as well as the cone angle ranges of the N = 11 and 13 clusters. A faster cooling rate, however, encourages the existence of metastable structures. The suppressing effect of a slow cooling rate suggests that these clusters are actually stable intermediates, or metastable.

#### 3.5. Cluster Structures at Larger N and the Implication for Previous Theoretical Studies

In contrast to uniquely packed polyhedral clusters at small to moderate N, hard cone-shaped particles form spherical clusters without well-defined local packings at larger N (or at small cone angles). This finding is consistent with a previous theoretical prediction\textsuperscript{11} and previous simulation results,\textsuperscript{13} as shown in Figure 6, where two larger hollow spherical clusters are obtained with 206 and 475 cone-shaped particles, respectively.

Some possible explanations for this observation are as follows. For hard cones, a spherical cluster has a higher packing fraction, or a lower potential energy, and thus is more stable or robust than a cylindrical cluster.\textsuperscript{11} Additionally, a spherical cluster has a higher symmetry and thus a larger configurational entropy, which may also contribute to the persistently observed spherical clusters in our cone-shaped particle simulations. As for the local packing, according to Euler’s theorem,\textsuperscript{26} 12 disclinations, i.e., 12 fivefold vertices, are required to close a hexagonal network, and different ways of distributing the 12 disclinations dictate the different local packings inside the spherical cluster, resulting in the disclinations on the surface of the spherical clusters in Figure 6.

The Israelachvili packing rule is often used to rationalize the structures assembled from surfactants and amphiphilic particles. It is worth noting that no cylindrical clusters are observed in our simulation, in contrast to the prediction of the Israelachvili rule for truncated cones. Our observation, however, may be limited by the following: (1) We investigated rigid cone systems, while the Israelachvili packing rule was initially conceived for soft amphiphiles. (2) Our model contains attractive interactions between interior beads only, and thus, the cones are not simply analogous to rigid bulky surfactants. (3) Our simulations are all carried out in the dilute regime where clusters are favored.

Iacovella et al.\textsuperscript{27} applied the Israelachvili packing factor\textsuperscript{10} to a monotethered nanosphere system. They calculated the shape factor from the effective length of the tail (or tether), the effective volume of the tail, and the effective area of the head group (or nanoparticle). For a fixed building block concentration and head diameter, they found transitions from spherical micelles to hexagonally packed cylinders to lamellar bilayers with increasing tether length, which is consistent with the theoretical predictions of Israelachvili. This indicates that the Israelachvili packing rule is more suitable for describing the behavior of amphiphiles of soft conical particles.

A similar situation was also observed by Park et al.,\textsuperscript{3} where rodlike metal–polymer amphiphilic particles were considered as truncated cones with soft, aggregating tails and were qualitatively analyzed using the Israelachvili rule to explain the appearance of self-assembled tubular and sheet structures. Modeling the peptide amphiphile as a lower, rigid part and an upper, flexible part and using molecular dynamics in 2D space, Tsonchev et al.\textsuperscript{28} found that long-range dipole interactions in the flexible part and hydrogen bonding in the rigid parts together stabilize peptide amphiphile self-assembly into cylindrical micelles as found in experiments.\textsuperscript{2} As such, a model of soft cones with long-range interactions can be obtained by suitable modification to the current model.

4. Conclusions

In summary, we investigated the self-assembly of anisotropic rigid cone-shape-shaped particles with directional attractions and demonstrated that structures with pre-designed geometric accuracy could be obtained by the self-assembly of such building blocks. We presented a diagram that can be referenced to predict the self-assembled structure for a given cone angle, providing important guidance for the design and assembly of conical building blocks. Specifically, we found that when $N$, the number of cone-shaped particles in a cluster, is small, the finite packings obtained from the self-assembly process produce a series of distinct cluster structures that resemble the packings found in evaporation-driven assembly of colloidal spheres. When $N$ is large, spherelike clusters, instead of cylindrical clusters as predicted by the Israelachvili rule for truncated cones, are always found, which is consistent with previous predictions by the geometric packing analysis and MD simulations on hard cones.

Moreover, we find that certain cone angle ranges have narrow cluster size distributions, indicating desirable stability and reproducibility, which suggests the superiority of conical building blocks over isotropic building blocks for the fabrication of precise structures. We classify our simulated precise structures into stable and metastable clusters by their wide and narrow cone angle ranges and influence of cooling rate, and by their monodisperse and nonmonodisperse cluster size distribution profiles and relatively high and low yields.

Additionally, we find that, for stable clusters, varying the cooling rate has little or no influence on their cone angle ranges and size distribution profiles. However, slower cooling rates can change cluster size distribution profiles and cluster yields significantly for metastable clusters. Furthermore, a slow cooling rate promotes thermodynamically stable clusters and suppresses metastable clusters, as expected.

The design principles identified in this work by investigating the dependence of the cluster packings and cluster size distribution on the cone angles enable programmable and predictable assembly of precise structures with patterned hard cones and may provide promising routes of fabricating novel designer materials. While the specific angle ranges of stability will undoubtedly change somewhat for cones of different sizes or different interaction ranges, the diagram provides an accurate prediction for cones matching the specifications modeled here and general design principles for cones of different specifications. The better understanding achieved in this work on the assembly of anisotropic conical particles may also help explain and control the structure of matter at different length scales for future applications.

Finally, how might such ringlike attractive patches in our model cone be introduced onto cone-shaped particles in experiments? Several experimental approaches appear promising. Sheu et al.\textsuperscript{6,7} obtained uniform nonspherical particles such as ice-cream-cone-like particles from identical polystyrene polymers. It may be possible to extend this technique to synthesize cone-like particles from different polymers and convey different properties to the cap and lower part of the ice-cream-cone-shaped particle, for example, by making the cap hydrophilic and the lower part hydrophobic. The hydrophobic interaction with solvent may thus drive the assembly of particles into structures such as those studied here. Alternatively, a decorating technique newly developed by Stellacci and co-workers\textsuperscript{29} may also be used where ordered phase-separated domains of stabilizing ligands spontaneously occur on the curved surface of nanoparticles.

It is worth mentioning that the model and simulation approach described in this work can be easily extended to study other anisotropic colloidal particles that have been recently synthesized, such as dumbbell-like,\textsuperscript{6,7,30} peanut-like,\textsuperscript{30,31} ellipsoid-like,\textsuperscript{32} and spindle-like\textsuperscript{30} particles.

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