Self-assembly of monodisperse clusters: Dependence on target geometry

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We apply a simple model system of patchy particles to study monodisperse self-assembly, using the Platonic solids as target structures. We find marked differences between the assembly behaviours of the different systems. Tetrahedra, octahedra and icosahedra assemble easily, while cubes do not. We relate these differences to the kinetics and thermodynamics of assembly, with the formation of large disordered aggregates a particular important competitor to correct assembly. In particular, the free energy landscapes of those targets that are easy to assemble are funnel-like, whereas for the dodecahedral system the landscape is relatively flat with little driving force to facilitate escape from disordered aggregates.

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I. INTRODUCTION

The assembly of nano-structured materials and devices presents a great challenge for the future. A very promising approach is offered by self-assembly processes, in which nanoscopic or colloidal building blocks come together spontaneously to form ordered structures. While top-down approaches to assembly become increasingly challenging on small length scales, self-assembly offers a bottom-up approach which circumvents many of these difficulties, while providing its own unique challenges in the synthesis and design of the subunits.

Current examples of synthetic self-assembling systems, such as micelles and block copolymers, usually lead to polydisperse and imperfectly controlled products. Biological systems, by contrast, display an astonishing variety of ordered and precise self-assembly processes1 For example, virus replication involves the assembly of hundreds of proteins to form highly symmetric and monodisperse shells (capsids). Such systems provide an inspiring example of the level of control possible in self-assembly. Future applications in nanotechnology are likely to require this level of sophisticated control in order to form precisely ordered structures.

One attractive aspect of self-assembly is the idea that the structure of the final product is entirely determined by the interactions between the subunits (along with the reaction conditions). Hence, given sufficient understanding of the principles of self-assembly, the subunits can be designed to produce a given product. In order to implement these designs a great deal of control is required over the structure of the subunits. One key requirement for sophisticated self-assembly is the production of anisotropic subunits, which only attract each other in certain directions. Rational design of this anisotropy, both in the shape and in the interactions, then allows for control over the resultant self-assembled structures.

Recently there has been a great deal of progress towards synthesising such anisotropic subunits2,3,4. Promising avenues included nanoparticles covered with mixed thiol monolayers,5,6 two-faced Janus,7,8 and even triphasic9 particles created at interfaces, small colloidal clusters,10,11 where the gaps between the individual colloid can be filled with a controlled amount of another material to create symmetrical ‘patchy’ particles, and colloids with patches grown at the sites of contacts in a crystal.12

The production of functional self-assembling systems will require not only advanced synthesis techniques to produce the subunits, but also a strong theoretical understanding of the self-assembly process so that appropriate designs and conditions can be chosen. Recently there has been significant progress in the theory13–20 and simulation21–34 of monodisperse self-assembly. However, these studies have mostly focussed on the self-assembly of virus capsids or other protein complexes, and so include protein-like interactions which are more specific than those which are likely to be available in the first generation of synthetic patchy particles.

Here we consider a minimal patchy particle model, representing spherical colloids or nanoparticles with anisotropic interactions. Our model is intentionally simple, so that we are easily able to observe and study behaviours which we expect to be general properties of monodisperse self-assembly rather than being specific to our model. This simplicity also allows us to comprehensively survey the behaviour, and to connect the kinetics with the form of the underlying free energy landscapes.

We have previously used the model to study the self-assembly of 12-particle icosahedra.35 Here, we build on this work to consider the effect of the target geometry on the self-assembly process, using the Platonic solids as our targets. We use these examples to address questions such as whether self-assembly becomes more difficult with increasing target size, and whether particular geometric features make some targets particularly difficult to as-
semble. In doing so, we will pay particular attention to the main processes which compete with successful assembly, and how their impact on the yield can be minimised. We hope that the design principles we learn from this work will offer some guidance to experimental groups seeking to make practical synthetic self-assembling systems.

Although the assembly process studied here is similar to that of virus capsids, in that anisotropic particles come together to form closed, highly symmetric shell structures, the interaction potential we use has no dependence on the torsional angle between interacting particles. While this is likely to be a good choice for modelling synthetic anisotropic particles, it is not a good representation of the interactions between proteins, and leads to behaviour not observed for systems of virus capsomers. We consider a model including torsional interactions, and leads to the main processes which compete with successful assembly, while allowing for efficient simulation. The model consists of spherical particles patterned with attractive patches. They are described by a modified Lennard-Jones potential, in which the repulsive part of the potential is isotropic, but the attractive part is anisotropic and depends on the alignment of patches on interacting particles. Specifically, the potential is described by

$$V_{ij}(r_{ij}, \Omega_i, \Omega_j) = \begin{cases} V_{LJ}(r_{ij}) & r < \sigma_{LJ} \\ V_{LJ}(r_{ij}) V_{\text{ang}}(\hat{r}_{ij}, \Omega_i, \Omega_j) & r \geq \sigma_{LJ}, \end{cases}$$

where $V_{LJ}$, the Lennard-Jones potential, is given by

$$V_{LJ}(r) = 4\epsilon \left( \frac{\sigma_{LJ}}{r} \right)^{12} - \left( \frac{\sigma_{LJ}}{r} \right)^6.$$  \hspace{1cm} (2)

$V_{\text{ang}}$ is an angular modulation factor, which depends on the orientations of the patches on the two interacting particles, as well as the direction of the vector joining them. Specifically,

$$V_{\text{ang}}(\hat{r}_{ij}, \Omega_i, \Omega_j) = G_{ij}(\hat{r}_{ij}, \Omega_i) G_{ji}(\hat{r}_{ji}, \Omega_j),$$  \hspace{1cm} (3)

where

$$G_{ij}(\hat{r}_{ij}, \Omega_i) = \exp \left( -\frac{\theta_{k_{ij}}^2}{2\sigma^2} \right),$$  \hspace{1cm} (4)

$\sigma$ gives the width of the Gaussian, $\theta_{k_{ij}}$ is the angle between patch vector $k$ on particle $i$ and the interparticle vector $r_{ij}$, and $k_{\text{min}}$ is the patch that minimizes the magnitude of this angle. Hence, only the patches on each particle that are closest to the interparticle axis interact with each other, and $V_{\text{ang}}$ is one if the patches point directly at each other. One feature of this potential is that as $\sigma \to \infty$ the isotropic Lennard-Jones potential is recovered. For computational efficiency the potential is truncated and shifted at $r = 3 \sigma_{LJ}$, and the crossover distance in Eq. (1) is adjusted so that it still occurs when the potential is zero.

A particular particle is specified by a set of unit vectors describing the positions of the attractive patches. For each of our target structures, the patches are placed such that they point directly at the neighbouring particles in the target structure. Fig. 1 shows the component particles and complete clusters for each of our target structures, the Platonic solids. Note that for these targets all the particles and patches are equivalent.

Somewhat similar patchy particle models have been used to study gel formation and the crystallization of proteins and patchy colloids and fibre formation.

**II. METHODS**

**A. Model**

We make use of a minimal model, designed to contain only the essential features required for targeted self-assembly, while allowing for efficient simulation. The model consists of spherical particles patterned with attractive patches. They are described by a modified Lennard-Jones potential, in which the repulsive part of the potential is isotropic, but the attractive part is anisotropic and depends on the alignment of patches on interacting particles. Specifically, the potential is described by

$$V_{ij}(r_{ij}, \Omega_i, \Omega_j) = \begin{cases} V_{LJ}(r_{ij}) & r < \sigma_{LJ} \\ V_{LJ}(r_{ij}) V_{\text{ang}}(\hat{r}_{ij}, \Omega_i, \Omega_j) & r \geq \sigma_{LJ}, \end{cases}$$

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where

$$G_{ij}(\hat{r}_{ij}, \Omega_i) = \exp \left( -\frac{\theta_{k_{ij}}^2}{2\sigma^2} \right),$$  \hspace{1cm} (4)

$\sigma$ gives the width of the Gaussian, $\theta_{k_{ij}}$ is the angle between patch vector $k$ on particle $i$ and the interparticle

**B. Dynamical simulations**

In the simulations of our model we wish to represent the Brownian motion that colloids and nanoparticles undergo in solution. As we do not include any solvent particles in our coarse-grained description of the system, a simple and efficient way to represent this dynamics is to use Monte Carlo (MC) where the moves are restricted to be local, since this ensures that the dynamics are diffusive.

In particular, we use Metropolis MC in the canonical ensemble, using periodic boundary conditions. The allowed move types are small single-particle translations and rotations. The translational moves are randomly chosen from a cube centred on the selected particle. Rotational moves make use of a quaternion description of the particle’s orientation; the proposed quaternion is given by the renormalized sum of the current quaternion and a smaller, randomly generated 4-vector.

One potential problem with using single-particle moves is that, although free particles and clusters undergo diffu-
sion as required, the relative diffusion rates of clusters of different sizes can be incorrect with the larger clusters diffusing too slowly. However, in practice for systems where the main mechanism of cluster growth is by monomer addition rather than cluster-cluster aggregation, single-particle moves are sufficient. Indeed, preliminary simulations using the virtual move MC algorithm, which has been very recently introduced by Whitelam and Geissler and is designed to overcome this problem by using cluster moves,

show only very minor differences to those presented here. By contrast, we have found that such an algorithm is crucial for systems designed to assemble hierarchically.

\section*{C. Equilibrium simulations}

To compute free energy landscapes and the positions of thermodynamic transitions we make use of umbrella sampling. The essential idea is to bias the system such that free energy barriers are easier to cross, thus allowing the system to explore configuration space much more quickly. Under this scheme, instead of choosing the acceptance probabilities using a Boltzmann distribution, we use the modified distribution \( \exp [ -\beta (V + w(Q)) ] \), where \( Q \) is an order parameter and \( w(Q) \) is a weighting function. Canonical averages are simple to obtain from a simulation of such a non-Boltzmann \((nB)\) ensemble using the expression

\[
\langle B \rangle_{NVT} = \langle B \exp [\beta w(Q)] \rangle_{nB},
\]

where \( B \) is some generic property of the system.

Despite considerable effort, it did not prove feasible to devise an order parameter that facilitated the formation of multiple copies of the target structure. Therefore, we instead restricted our umbrella sampling simulations to the formation of a single target structure at the same density. There are, however, two possible reasons why the equilibrium data for this finite-sized system can differ from that for bulk. Firstly, it does not allow for interactions between the target clusters. But since the attractions between these clusters are small, this is a good approximation. Secondly, it does not allow for configurations where the fraction of the atoms in the target structure is non-integer; in our umbrella sampling simulations the only possibilities are for all the particles to be in the complete target structure or none of them. For example, at the midpoint of the transition, the small system will fluctuate in time between the target cluster and the monomeric gas state, spending half the time in each state, whereas for a large system one would expect that at any given time the configuration will be a mixture of target clusters and monomers with half the particles in the target clusters. Analytical calculations have indicated that this restriction has only a relatively small effect on the position of the centre of the transition between monomers and the target structure (this is what we are most interested in here), but can cause an appreciable narrowing of the transition compared to the thermodynamic limit.

A convenient order parameter to study the transition between a monomeric gas and the target structure is then the number of particles in the largest cluster in the simulation. The weighting function \( w(Q) \) was found by iteratively performing simulations and improving \( w(Q) \) at each iteration until the time spent at each value of \( Q \) was approximately equal.

In the case of the dodecahedron we found that the above order parameter was inadequate because the larger clusters that formed were disordered rather than based on the dodecahedron. We therefore introduced a second-order parameter, namely the number of pentagons in the largest cluster, with the aim of promoting the formation of dodecahedral clusters. However, this was still insufficient to drive the system to assemble successfully.

Instead, to finally achieve equilibrium we had to combine this two-dimensional umbrella-sampling scheme with Hamiltonian exchange. In the latter, additional exchange moves are introduced that involve swapping configurations between simulations with different Hamiltonians, the purpose being to couple the system of interest to one where equilibrium is easier to achieve. In our case, we know that successful assembly of dodecahedra is possible for a similar patchy particle model, but where a torsional component is included in the potential. Specifically, the attractive region of the potential is modulated by an additional factor

\[
V_{tor} = \exp \left( -\frac{\phi}{2\sigma_{tor}^2} \right),
\]

where \( \phi \) is a torsional angle between two particles, chosen such that \( \phi = 0 \) in the target structure. In the limit that \( \sigma_{tor} \) approaches infinity, our original potential is recovered.

In our Hamiltonian exchange scheme we ran an array of 20 simulations at different values of \( \sigma_{tor} \) ranging from 1.25 radians to \( \infty \). The intermediate values were chosen to maximise the rate of configurational exchange. The probability of an exchange move was 0.1 and only exchanges between adjacent values of \( \sigma_{tor} \) were attempted. The acceptance probability that ensures detailed balance is

\[
p_{acc}(i, 1, (j, 2) \rightarrow (i, 2), (j, 1)) = \min \{ 1, \exp [ -\beta (W_{i,1} + W_{j,2} - W_{i,2} - W_{j,1}) ] \}
\]

where \( i \) and \( j \) represent different Hamiltonians, and 1 and 2 different configurations. \( W_{i,1} \) is the combined potential energy and umbrella weighting of configuration 1 with the Hamiltonian \( i \), i.e. \( W_{i,1} = V_i(r_1^N) + w_i(Q(r_1^N)) \).

\section*{D. Structural Analysis}

Many of our results make use of information on the clusters present in the simulations, and of the rings
present within those clusters. We define two particles as being bonded if their interaction energy satisfies $V_{ij} \leq -0.4$. Particles are part of the same cluster if they are connected by an unbroken chain of bonds. We identify target clusters by matching the number of particles and bonds in the cluster to the profile for the target cluster, including some allowance for bonds temporarily weakened by thermal fluctuation. We define rings using the shortest-path algorithm given in Ref. 50; a closed loop is counted as a ring if there is no pair of particles on the ring connected by any chain of bonds shorter (i.e. consisting of fewer bonds) than the route by which they are connected along the ring.

III. RESULTS

We first wish to identify the conditions under which successful assembly of the target structure occurs. Because of the relative simplicity of our model, we are able to perform large numbers of simulations, and hence comprehensively map out the assembly behaviour over a range of parameter space. Fig. 2 shows the dependence of the final yields of each of the target structures on temperature (relative to $\epsilon$) and patch width in simulations starting from an initial random geometry after a given number of MC cycles. Although the yields in certain regions will tend to increase slightly with longer simulation times, the form of the diagrams is robust. Also note that low (high) temperature is equivalent to strong (weak) interactions.

To first order the general shapes of the plots are similar, because the same physical principles apply in each case. However, there are also significant differences. Tetrahedra, octahedra and icosahedra assemble readily over wide ranges of parameter space. By contrast, cubes are more difficult to assemble and are only formed in a smaller region of parameter space, while dodecahedra never form at all. We shall examine the reasons for these differences in detail later.

The region of successful assembly is determined by a number of constraints, both thermodynamic and kinetic. Firstly, we find that at high temperature $T$ the stable state becomes a gas of monomers and small clusters, and target clusters are no longer formed. A second thermodynamic constraint arises at high patch width $\sigma$, where the patches are so wide that the target clusters cease to be the most stable state of the system. Each patch then becomes capable of interacting with more than one neighbour, and so the system can both lower its energy and raise its entropy by forming large, unstructured liquid-like droplets.

Both of these effects can be seen in Fig. 3 which shows the average cluster size for each target structure, again as a function of $T$ and $\sigma$. At high $T$ the average cluster size approaches unity, signifying a monomer gas, while at high $\sigma$ the existence of a liquid phase results in clusters containing essentially all the particles in the simulation.

FIG. 2: (Colour Online) The percentage yield of target clusters formed after 80000 MC cycles as a function of the patch width $\sigma$ (measured in radians) and the temperature for 1200 particles at a number density of $0.15 \sigma^{-3}$, for particles designed to form (a) tetrahedra, (b) octahedra, (c) cubes and (d) icosahedra. No plot is included for particles designed to form dodecahedra, since under no conditions did any dodecahedra ever assemble. The white lines show the thermodynamic transition temperature $T_c$ for the transition from a gas of clusters to a gas of monomers calculated using umbrella sampling.
FIG. 3: (Colour Online) The mean cluster size (averaged over particles) of systems of particles designed to form (a) tetrahedra, (b) octahedra, (c) cubes, (d) icosahedra, and (e) dodecahedra, for the same simulations as Fig. 2. The white lines show $T_c$.

FIG. 4: A schematic diagram illustrating the regions of parameter space in which different behaviours are observed. $T_c$ is the midpoint of the transition between the cluster gas and the monomer gas. $T_{lv}$ and $T_{lc}$ are the liquid-vapour phase transition lines for the monomer and cluster gases, respectively. $T_{agg}$ is the temperature below which aggregates become unable to rearrange to form target clusters on the time scale of the simulations, instead becoming trapped in a glass-like state. $\sigma_k$ is the value of $\sigma$ below which the patches are so narrow that little assembly or aggregation is able to take place on the time scale of the simulations. $\sigma_{\text{min}}$ represents the lowest value of $\sigma$ at which the liquid state is stable and $\sigma_{\text{max}}$ the highest value of $\sigma$ at which the cluster phase is stable.

Kinetic constraints become important at low values of $T$ and $\sigma$. At low $T$ the system becomes unable to escape from incorrect configurations. Since it very likely that some incorrect bonds will be formed during the assembly process this results in very low yields, and the system instead forms glassy kinetic aggregates. At low values of $\sigma$ the patches are very narrow. Particles will only rarely be sufficiently well aligned to feel attractive interactions from their neighbours, and growth is suppressed.

Sandwiched in the middle of these regions of inhibited assembly is a region of parameter space in which the target is both thermodynamically stable and kinetically accessible. In this region good yields of the target structure are obtained. These effects are summarised in Fig. 4, which shows schematically the different regions of parameter space and the lines which separate them. We will now examine these features in more detail, focussing on their dependence on the target geometry.

A. The cluster-monomer gas transition

As there are only very weak attractive forces between clusters assembled into the target structures, $T_c$, the temperature above which the stable state is dominated by monomers rather than target clusters, represents the midpoint of a chemical equilibrium between a monomer gas and a cluster gas. The plots in Fig. 2 show that $T_c$
increases with patch width $\sigma$ for all of the systems. This feature can be explained by considering the free energies of the monomer and cluster gases. At $T_c$, $A_m$ and $A_c$, the free energies of the monomer and cluster gases respectively, are approximately equal. $A_m$ is largely independent of $\sigma$, as is the ground state potential energy per particle in the cluster gas, $V_c^{gs} \approx m\epsilon/2$, where $m$ is the number of patches per particle. However, the entropy $S_c$ of the cluster gas increases with $\sigma$. At higher $\sigma$ the clusters are free to undergo larger vibrations, leading to a higher vibrational entropy, and hence a larger $T_c$.

While this trend is a generic feature, $T_c$ also shows strong target dependence. Fig. 5 shows the heat capacity $C_v$ as a function of temperature for each of the target structures at $\sigma = 0.45$. The data was obtained from umbrella sampling simulations containing sufficient particles to form one target cluster.

Table I shows estimated values obtained using this equation, along with values obtained from umbrella sampling simulations. Despite the simplicity of the model, it matches the data reasonably well, and provides an explanation for the ordering of the $T_c$ values.

The value of $T_c$ has an important impact on the ability of a system to self-assemble, with lower values of $T_c$ making assembly more difficult, because the potential range of $T$ over which assembly might occur is decreased. It is noteworthy that cubes and dodecahedra have the lowest values of $T_c$, and are the hardest of our targets to assemble.

As the monomer-cluster transition is a chemical equilibrium, the heat capacity peaks have a finite width. The relative width is expected to decrease with increasing target size, and this can be seen in Fig. 5 with the exception of the dodecahedral system, again indicating that this system is behaving differently.

### B. The cluster-liquid transition

For all systems, we see the formation of large aggregates both at sufficiently large values of $\sigma$ and at low temperature for narrower patches (Fig. 3). The aggregates formed in these two regions are in fact closely related, with their properties varying smoothly between them, as is especially clear for the dodecahedral case, where there is no region of cluster formation separating the two regions of aggregate formation. Nevertheless, it is natural to divide these aggregates into two groups. In the first, at high $\sigma$, the aggregates are the stable state of the system. In general these thermodynamic aggregates have some liquid-like characteristics, being fairly compact and undergoing constant rearrangement. The second group of aggregates, those found at low temperature and lower $\sigma$, are metastable, with the target cluster being the true stable state. These kinetic aggregates also display more gel-like properties, often forming extended networks of string-like formations, which may percolate. However, we have not sought to establish whether they form as a result of kinetically-arrested liquid-vapour phase separation, or gelation of a homogeneous fluid.

The aggregates tend to have irregular forms. In the
case of the thermodynamic aggregates, so long as all patches are pointing inwards, the surface tension is relatively low, because the energetic cost of the surface is small. As a result there is a significantly smaller driving force for forming spherical droplets than for systems with isotropic interactions. In the case of the kinetic aggregates, the reduced ability to rearrange because of insufficient thermal energy and narrow patches leads to the formation of increasingly irregular and ramified aggregates at low $\sigma$. The energy of the aggregates tends to become less negative as well, since these increasingly glassy systems are less able to optimise their configurations to achieve lower energies. This effect is clearly visible in Fig. 6, which shows the dependence of the final energies in the simulations on $T$ and $\sigma$ for each of the target structures.

The structures of the aggregates also have a strong dependence on the target. At temperatures and patch widths close to those where target clusters can successfully form, the local structure within the aggregates tends to have a strong similarity to that in the target cluster, so that icosahedron-like groups are visible in the aggregates of icosahedron-forming particles, and so on. These structural similarities are visible in Fig. 7 (a)–(d), which shows snapshots of liquid aggregates near to the region of successful assembly. The one target which consistently produces aggregates with little resemblance to the target structure is the dodecahedron. The wide splay angle (i.e. the angle between the patches and the symmetry axis) allows the formation of a wide variety of structures, resulting in highly ramified structures containing many hexagons and larger rings as well as pentagons (see Fig. 7(c)). However as we move to higher values of $\sigma$ the differences between the aggregates formed for the different targets are lost, so that the dense aggregates formed at very high $\sigma$ look essentially the same for all cases.

Fig. 7(f)–(j) show typical kinetic aggregate structures. We can see that all of the kinetic aggregates have a tendency to form chain-like structures. In larger simulations these may sometimes connect together and percolate across the simulation box. The local structure of the chains is again heavily dependent on the target structure, reflecting the symmetries of the target. We can also see that the tetrahedra have a greater tendency to form a number of smaller aggregates, which is consistent with Fig. 3. The narrow splay angle of the tetrahedron-forming particles promotes high curvature and hence small clusters. This effect is lost at higher $\sigma$, as the patch positions begin to have less control over aggregate structure.

Interestingly, similar ramified worm-like aggregates have also been seen for Janus particles that have a hydrophobic and a charged side. The similarity is probably because of the effectively one-sided nature of the attractions for our patchy particles.

The extent of the regions in which aggregates are found is important, as it sets boundaries on the region in which the target structure may successfully be formed. In this

FIG. 6: (Colour Online) The normalised final total energy $E/2Nm$ for systems of particles designed to form (a) tetra- hedra, (b) octahedra, (c) cubes, (d) icosahedra, and (e) dodecahedra, for the same simulations as Fig. 2. The energy is normalised such that a value of $-1$ indicates that, on average, all patches are involved in one perfect bonding interaction. The white lines again show $T_c$. 


section we consider the factors affecting the region in which liquid-like aggregates are stable, while in Section III C we will return to consider those determining the extent of the kinetic aggregate region.

It is useful to define some terms to describe the position of the transition from the cluster phase to the liquid at high $\sigma$. We define $\sigma_c^{\text{max}}$ as the maximum value of $\sigma$ at which the cluster phase is found (which will be at zero temperature) and $\sigma_c^{\text{min}}$ as the minimum value of $\sigma$ at which the liquid phase is found. These two points are connected by the transition line $T_{lc}$. All three are shown in Fig. 4. $\sigma_c^{\text{max}}$ depends entirely on the energies of the two states, while the gradient of $T_{lc}$ depends on the entropy difference between the two states.

Fig. 5 shows that the dodecahedral and cube-forming particles are the most able to satisfy their patchy interactions in liquid droplets, and attain the greatest bonding energies. The low number of patches per particle $m$ and the wide angles between patches result in fewer constraints on the liquid structure, making it easier to optimise patch-patch interactions. We can see a clear correlation between the angles between patches and the energy of the liquid. For the tetrahedral, octahedral and icosahedral particles, with angles between the patches of 60°, the particles would be required to pack very closely together in order to satisfy all their bonds, and in fact we see relatively small bonding energies per patch. As we increase the angle, to 90° for cubes and then 108° for dodecahedra, the energy per patch steadily increases in magnitude.

The entropy of the aggregates might be expected to be correlated with the energy. When there are fewer constraints it is not only easier to satisfy patch-patch interactions in a disordered configuration, but there are more ways of doing so, leading to a higher configurational entropy. We would therefore expect larger aggregate entropies for the dodecahedral- and cube-forming particles.

We can obtain some information about the relative entropies of the aggregate and cluster phases by considering $T_{lc}$. The shallower the gradient of $T_{lc}$ the greater the entropy differences between the aggregate and cluster phases, with negative gradients indicating that the aggregates are entropically favoured. Although calculating precise values of $T_{lc}$ would be far from straightforward, we can get an impression of how $T_{lc}$ varies with $\sigma$ from Fig. 3. From those plots we can see that $T_{lc}$ has a negative gradient for octahedral, icosahedral and probably cubes, indicating that the liquid has a higher entropy than the cluster gas, and hence that the disorder in the liquid more than compensates for the loss of translational degrees of freedom on forming a large aggregate. $T_{lc}$ is close to vertical for the tetrahedral system indicating that the entropies for the liquid and cluster gas are similar—as the smallest of the targets, the tetrahedral clusters have the largest translational entropy. The gradient for the dodecahedral systems is unclear from Fig. 3 because of the absence of target formation. Later in Sec. III D we will see that $T_{lc}$ has a far shallower slope for dodecahedra, which is consistent with our expectations based on the low energy of the aggregates and their large cluster size.

C. Mechanisms of assembly and misassembly

The dynamics of the self-assembly simulations depend strongly on temperature and on the target. Fig. 8 shows the yields of each of the target structures as a function of $T$ and time at $\sigma = 0.45$. These plots show a number

![Colour Online) Typical aggregate configurations (a)–(e) in the liquid-vapour coexistence region but (a)–(d) close to the region of successful assembly, and (f)–(j) in the kinetic aggregation regime for particles designed to form (a) and (f) tetrahedra, (b) and (g) octahedra, (c) and (h) cubes, (d) and (i) icosahedra, and (e) and (j) dodecahedra. The values of $(kT/\epsilon, \sigma)$ are (a) (0.06, 0.65), (b) (0.09, 0.65), (c) (0.08, 0.55), (d) (0.16, 0.6), (e) (0.04, 0.6), (f) (0.05, 0.35), (g) (0.07, 0.35), (h) (0.06, 0.4), (i) (0.08, 0.35) and (j) (0.04, 0.35). Each simulation contains 60 particles at a number density of 0.15 $\sigma_{LJ}^{-3}$.](image)
of interesting features.

Firstly, in each case the yield approaches equilibrium most rapidly at high temperatures, close to $T_c$. Here the dynamics are relatively fast, and since liquid-like aggregates are not stable with respect to the monomer gas at these temperatures, the clusters face no competition. However, close to $T_c$ the yield is limited by the finite width of the cluster gas-monomer gas transition, with smaller clusters displaying a broader transition, as is clear in Fig. [5]

At slightly lower temperatures the time taken to reach equilibrium becomes longer, both because the time scale for rearrangement and the breaking of incorrect bonds is longer and because the diffusional time scales increase as the assembly progresses towards higher yields. As a result the temperature at which optimal yields are obtained decreases with time, being determined by the competition between rapid assembly at high temperatures and higher equilibrium yields at lower temperatures.

Moving to still lower temperatures the yield on the time scale of our simulations falls off, until around $T_{agg}$ it becomes severely limited. For target structures of tetrahedra, cubes and icosahedra, reasonably sharp cut-offs are seen at low $T$, giving approximate values for $T_{agg}$. In the case of the tetrahedra the yield does not fall to zero but rather to a finite value of around 20%, as some yield of tetrahedra is expected by straightforward chance assembly with no rearrangement required. Unlike the other targets, the plots for octahedra show an approximately linear decay in yields with decreasing temperature. The reasons for this are unclear.

Fig. [8(a)] shows the yield as a function of time for each target structure (except the dodecahedron) at the optimal conditions for the formation of that structure in Fig. [2] (i.e. those that give the maximum yield after 80000 cycles). While the final yields obtained in each case approach 100%, the time taken varies by almost two orders of magnitude. The order of the time taken for tetrahedra, octahedra and icosahedra is consistent with their relative sizes, with larger clusters taking longer to assemble as expected. Some tetrahedra form at very short times via the chance assembly mentioned above. However, the cubes take anomalously long to assemble, which is partly explained by the narrow range of temperature over which cubes can assemble (Fig. [8(c)]) because of the low value of $T_c$ for this system.

We will now consider in more detail the dynamics in the region of parameter space where the target clusters are most stable by considering three sub-regions of that space. Firstly, there is the regime where the target cluster is the only structure stable with respect to the monomer gas (i.e. $T_{lv} < T < T_c$). As a result assembly occurs by direct nucleation of the target structure by the addition of monomers and other small clusters to a growing cluster, rather than proceeding via an aggregate state. Because aggregates are not stable there are essentially no competing states to the target structure.

Note that as this region is close to $T_c$, the yield may be

![FIG. 8: (Colour Online) The yields of (a) tetrahedra, (b) octahedra, (c) cubes, and (d) icosahedra after different numbers of simulation steps as a function of temperature $T$, in simulations of systems of 1200 particles at $\sigma = 0.45$ and at a number density of $0.15\,\sigma_{LJ}^{-3}$. Each data point is an average from five simulations.](image_url)
In the second regime both the target clusters and large liquid aggregates are stable with respect to the monomer gas (i.e. $T < T_c$ and $T < T_{1w}$ but $T > T_{agg}^{ss}$), leading to competition. However, mobility in the liquid droplets allows them to rearrange to form target clusters. Once complete clusters are formed all their patches point inwards, such that they experience very little attractive interaction with other particles. They are therefore able to “bud off”, separating from the rest of the liquid droplet in which they formed.

The optimal conditions for assembly are typically found close to the boundary of these two regimes, at a point in parameter space where both the direct nucleation and “budding-off” mechanisms of assembly operate. Fig. 9(b) shows the average cluster size against time under optimal assembly conditions for each of the target structures. For the cubes (and to some extent the octahedra) the average size passes through a maximum before falling to the target cluster size, indicating clearly that the mechanism of aggregation followed by rearrangement plays a significant role. Even though the “budding off” mechanism is not sufficiently prominent to cause a maximum in the cluster size for tetrahedra and icosahedra, it still contributes to the rapidity of assembly under these conditions.

For $T < T_{agg}^{ss}$ assembly is suppressed because the aggregates act as kinetic traps. Hence, we define $T_{agg}^{ss}$ as the temperature below which large clusters have insufficient thermal energy to rearrange to form target clusters on the time scale of the simulation (and is therefore weakly dependent on time scale). Below $T_{agg}^{ss}$ generally only large aggregates are formed, often forming extended ramified networks which may percolate.

$T_{agg}^{ss}$ is dependent on target geometry for a number of reasons. The first factor is the degree of order in the aggregates. Fig. 7 shows that the aggregates can contain a considerable degree of local structure that is similar to the target, and hence only a relatively small amount of rearrangement is required for the aggregates to form complete clusters. The exception is the dodecahedral case, where the aggregates have very little dodecahedron-like structure at all, and would require almost total rearrangement to form the target clusters.

A second factor is the stability of the aggregates, because the more stable they are, the smaller is the thermodynamic driving force for them to rearrange to find the target structure. This factor is particularly relevant to the cubic and dodecahedral systems, which we saw earlier, had particular low energies and high entropies.

A third factor is the size of the aggregates. In general a smaller splay angle in the assembling particles tends to lead to a higher surface curvature and hence to smaller aggregates, which are easier to rearrange. In Fig. 8 we can see that the low-$T$ aggregates formed with tetrahedra and octahedra as target structures are much smaller than for the other shapes.

A final factor is the size of the target structures themselves. Less rearrangement is needed to form a smaller target. Further, for smaller targets, the growth of aggregates is less likely to deviate significantly from the correct assembly pathway. For example, for tetrahedra, some degree of successful assembly is observed even at very low $T$, because a significant proportion tetramers will have come together by chance to directly form a tetrahedron without the need for any rearrangement. However, even for octahedra this effect becomes negligible.
D. The difficulty of assembling dodecahedra

Of the target structures examined in this paper, the dodecahedron is exceptional in the difficulty of its assembly. While the other challenging target, the cube, shows a restricted region in which assembly occurs, it is nevertheless easy to obtain high yields by conducting long simulations with carefully chosen parameters. By contrast dodecahedra are never formed with our model potential under any set of conditions.

In order to more fully understand the difficulty of assembling dodecahedra we attempted to obtain equilibrium thermodynamic data for a system of 20 particles, sufficient to assemble one dodecahedron. This was achieved only with considerable difficulty. Indeed, the pathological difficulty of forming a single dodecahedron even using biasing techniques underlines the extreme unlikeliness of ever forming one under ordinary dynamics, and implies the presence of a large kinetic barrier to assembly. Success was eventually achieved by combining two-dimensional umbrella sampling (using the number of bonds and the number of pentagons in the largest cluster as order parameters) with Hamiltonian exchange as described in Section II C.

In general, because of the low energy of the target structures and the specificity of the interactions, one would imagine that at moderate $\sigma$ there must exist a temperature range over which the target is the only stable structure with regard to the monomer gas, and so where aggregation formation does not compete with the assembly of the target. One might expect that in this region at least, dodecahedra should be able to assemble, so long as the time scales are sufficiently long that the nucleation barrier can be overcome. However, as we will see, this is in fact not the case, since over most of the range of $\sigma$ this region does not exist for the dodecahedral system.

Fig. 10(a) shows a plot of the heat capacity $C_v$ along with a plot of the average cluster size for $\sigma = 0.45$. At temperatures below the peak in $C_v$, the dodecahedron is most stable. However, we can see that the average cluster size remains high beyond this point, indicating that the dodecahedron first melts before gradually evaporating at higher temperature. The evaporation of this liquid cluster corresponds to the shoulder in the $C_v$ plot.

The generality of this picture is confirmed in Fig. 10(b). The higher line in the figure shows the temperature at which the average cluster size in the box has a value of 10.5 (the midpoint of 1 and 20), and the lower line $T_m$, the temperature at which a dodecahedron melts. The presence of an intermediate liquid state for this 20-particle system, which persists down to $\sigma \approx 0.25$, was a surprise to us. For, although we expected the free energies of the monomeric and cluster states to be relatively little affected by the small number of particles in these equilibrium simulations, we expected the liquid state to be considerably destabilized by the high surface to volume ratio of any 20-particle liquid cluster. That liquid clusters are seen in these simulations is testament to the low energy and high entropy of the aggregates for the dodecahedral system that we noted earlier. Thus, we expect the region of stability for the liquid state to be larger for a bulk system than for this 20-particle system, i.e. $T_{10.5}$ can be considered as a lower bound to $T_m$, $T_m$ as an upper bound to $T_c$, and 0.25 is an upper bound for $\sigma_{\text{min}}$

These results thus allow us to begin to understand why the dodecahedra are so hard to assemble. Given that at moderate $\sigma$ the stable state of the system at higher temperatures is as liquid droplets, the only region in which dodecahedra might be able to form is at lower temperatures, where, although they represent the global free energy minimum, aggregate formation competes with target assembly. In particular, in this region the system will aggregate very quickly, and so for a dodecahedron to form these aggregates would then need to be able to rearrange sufficiently, a process which is severely inhibited by the slow dynamics at these low temperatures. For
the dodecahedral system though, such rearrangement appears to be far more difficult than would be expected even allowing for temperature. This is probably a result of geometric factors which are unique, within our set of targets, to the dodecahedron. Because of the widely spaced patches on the particle surfaces, an enormous range of incorrect configurations are possible which satisfy the bonding well, and which deviate significantly from the target structure even on a local scale. For example, as well as pentagons, (non-planar) hexagons or larger polygons can form without introducing strain. Furthermore, it is possible for bonds to pass through these rings, leading to entangled networks of particles. These structures have little in common with correctly formed dodecahedra, and further, they experience little pressure to rearrange.

In order to obtain a clearer picture of the thermodynamic constraints on the dynamics of the system, we made use of data from the Hamiltonian exchange simulations to plot free energy landscapes in Fig. 11 and 12 not only for dodecahedra but also for icosahedra and cubes. The icosahedral system provides a contrasting example of a successfully assembling system. All are at a temperature corresponding to the maximum in the heat capacity.

**Fig. 11:** (Colour Online) Free energy landscapes for systems of (a) 12 icosahedron-forming particles, and (b) 20 dodecahedron-forming particles at $T = T_c$ and $\sigma = 0.45$. The order parameters are the numbers of particles and triangles/pentagons in the largest cluster.

**Fig. 12:** (Colour Online) Free energy landscapes for systems of (a) 12 icosahedron-forming particles, (b) 8 cube-forming particles and (c) 20 dodecahedron-forming particles at $T = T_c$ and $\sigma = 0.45$. The order parameters are the configurational energy and a measure of radial disorder, described in the text.

Fig. 11 shows the free energy as a function of the number of particles and the number of correct polygons (triangles for the icosahedron and pentagons for the dodecahedron) in the largest cluster in the system. In Fig. 11(a) we can see clear free energy minima representing the monomer gas and the complete icosahedron. Between the two minima is a region of higher free energy, with a free energy transition state with a significant amount of triangular ordering. Once the transition state is passed, the shape of the landscape directs the system towards forming an icosahedron, facilitating assembly. (Note that the high free energy of clusters of 12 particles with 19 triangles simply represents the fact that if one bond of an
icosahedron is broken two triangles are lost, and does not represent a barrier.)

By contrast, the monomeric gas is not a free energy minimum for the dodecahedral system at temperatures at which the dodecahedron is most stable (Fig. 11(b)). There is no barrier to the formation of a 20-particle cluster, so the system will tend to rapidly aggregate, and will then experience little drive to form a dodecahedron. Indeed, if the system follows the steepest downhill path it will tend to form a single aggregate containing few pentagons, so that significant rearrangement would be needed to form a dodecahedron.

Fig. 12 shows the free energy as a function of two different order parameters, the potential energy and a radial disorder parameter. The latter is defined as the standard deviation in the distance of particles from the centre of mass, and has a value of zero when all particles lie on a spherical shell (such as in the target structures). Again, the plot for the icosahedron shows two free energy minima, corresponding to a monomer gas and to an icosahedron, separated by a transition state of higher free energy. Importantly, we see that in order to obtain lower energies the system is forced to steadily reduce its maximum radial disorder, such that it is directed into the icosahedral state. The energy landscape is funnel-like.

By contrast, the free energy of the dodecahedral system appears to be almost entirely a function of the configurational energy, with very little dependence on the radial disorder. Energies close to that of the global minimum can be reached by highly disordered states. As a result the free energy landscape is relatively flat, and the system experiences very little driving force to form an ordered cluster. We also depict the landscape for cube-forming particles in Fig. 12(b). The intermediate difficulty of assembling the cubes is reflected in the broader, less-funneled nature of the landscape compared to the icosahedron, but there is still some free energy gradient towards the target structure.

Nevertheless, neither of the dodecahedral plots fully explain the situation. From these diagrams one might expect the formation of dodecahedra to be rare, but not essentially impossible. It appears that there is a large kinetic barrier preventing the formation of dodecahedra, which is not visible in these diagrams due to the choice of order parameters. Despite considerable efforts, we were not able to find a combination of order parameters which would show this barrier explicitly.

IV. CONCLUSIONS

We have presented a study of the self-assembly of a class of simple targets, the Platonic solids, using a minimal model of assembling patchy particles. We have comprehensively mapped out the behaviours of systems of particles designed to form each of the targets as a function of temperature and patch width. Further, we have obtained equilibrium data using umbrella sampling and Hamiltonian exchange, allowing us to compare the thermodynamic properties of the different systems, and to produce free energy landscapes for the larger targets to elucidate the differences in their behaviour.

We find that the behaviour of the systems and the success of self-assembly are strongly dependent on the target structure. One key property which varies considerably between targets is the stability of disordered aggregates, which have a complex relationship with the self-assembly process. Over large regions of parameter space they act as competition and prevent successful assembly, but in other regions assembly may proceed by first forming aggregates which then rearrange to “bud off” the target structures. As a result the assembly of cubes, whose constituent particles form particularly stable aggregates, is dominated by this budding mechanism, while dodecahedron-forming particles form aggregates so stable that they effectively block assembly of dodecahedra. In general we find that the stability of aggregates relative to the target clusters is determined largely by the spacing between the patches on the particles’ surfaces. Wider patch spacing leads to fewer constraints on the structures that the aggregates can adopt while satisfying all the patchy interactions, which in turn leads to both lower energies and higher entropies.

The stability of the target clusters themselves also varies considerably between targets, and depends both on the number of interactions each particle participates in, and on the size of the cluster, where greater size tends in general to decrease stability. High stability is a desirable trait as it allows assembly at high temperatures (or equivalently for weak interactions), where the breakup and rearrangement of misformed structures is more rapid.

We find that the dependence on shape can be such that certain shapes never assemble successfully; dodecahedra appear to be essentially impossible to correctly assemble in our model. The dodecahedron is a relatively unstable target because of its small number of interactions (each particle having only three nearest neighbours) and large size. The aggregates with which it competes, by contrast, are exceptionally stable. As a result at moderate patch widths the aggregates are stable to higher temperatures than the dodecahedra, and indeed there is no region of parameter space where the dodecahedron is both thermodynamically stable and kinetically accessible.

These observations collectively suggest a number of design rules for targets that will assemble easily. Firstly, a high number of nearest neighbours allows for more interaction which stabilises the target structure. Secondly, closely spaced patches serve to inhibit the formation of stable aggregates which can provide competition. These two features can be combined by choosing shapes with triangular faces, and as such we would argue that triangular faces are a major advantage in any target structure.

Proceeding to larger and more ambitious self-assembly targets, a difficulty arises. As the target size increases, the competition between the target and disordered aggregates increasingly favours aggregates, as the targets
become less entropically favourable. This suggests limits to the size and complexity of targets of monodisperse self-assembly which can be successfully formed using the simple interactions we have used here. One approach to circumvent this difficulty might be to use multi-component systems with different patch types that interact selectively, the latter being potentially achievable using DNA-mediated interactions between the particles. An alternative approach to this difficulty would be to introduce further constraints to the potential. In particular a potential including torsional constraints, i.e. one in which the interactions are specific both in direction and in relative orientation, would massively reduce the number of competing configurations available. Protein-protein interactions have this kind of specificity, and we expect that this is crucial in the assembly of large structures such as virus capsids. We examine the effect of using such a modified potential in the accompanying paper.

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