

View Article Online **PAPER** 



Cite this: Soft Matter, 2020, 16, 2774

Received 7th January 2020, Accepted 17th February 2020

DOI: 10.1039/d0sm00044b

rsc.li/soft-matter-journal

# Hierarchical self-assembly of patchy colloidal platelets†

Carina Karner, (1) \*\* Christoph Dellago and Emanuela Bianchi (1) bc

Anisotropy at the level of the inter-particle interaction provides the particles with specific instructions for the self-assembly of target structures. The ability to synthesize non-spherical colloids, together with the possibility of controlling the particle bonding pattern via suitably placed interaction sites, is nowadays enlarging the playing field for materials design. We consider a model of anisotropic colloidal platelets with regular rhombic shape and two attractive sites placed along adjacent edges and we run Monte Carlo simulations in two-dimensions to investigate the two-stage assembly of these units into clusters with well-defined symmetries and, subsequently, into extended lattices. Our focus is on how the site positioning and site-site attraction strength can be tuned to obtain micellar aggregates that are robust enough to successively undergo to a second-stage assembly from sparse clusters into a stable hexagonal lattice

## 1 Introduction

Nowadays, colloids of different shapes can be accurately synthesized within a vast range of symmetries, from complex convex units - such as spheroidal colloidal molecules, rods or polyhedra - to concave shapes - such as multi-pods or bowlshaped colloids. 1-3 On top of the shape-dependent anisotropic interaction, directionality in bonding is often sought after in order to fine tune the assembly of the particles into either finite clusters<sup>4-7</sup> or extended crystals.<sup>8-10</sup> The combination of bonding sites or extended surface patches<sup>11–13</sup> – i.e., regions of the particle surface where bonding occurs - and non-spherical particle shapes imparts preferred bonding directions between the particles, thus creating a rich enthalpic versus entropic competition to be taken advantage of for materials design.<sup>14</sup>

Finite clusters can serve, for instance, as prototypes of micro-robots to perform tasks at the micro-scale such as delivery and targeting. Recent and successful examples of anisotropic patchy particles assembling into finite clusters are colloidal asymmetric dumbbells that show a rotational propulsion under electric field, 15 dielectric cubes with one metallic facet that reconfigure on demand, 16 or Janus spheroids that act as encapsulating agents. 17

Self-assembled finite clusters can be also used as nonspherical building blocks for further - hierarchical - assembly

into extended structures and thereby expand the versatility of the original units. 18-22 Colloidal molecules composed of different types of particles, for instance, can be designed to support the assembly of superstructures with target photonic or phononic properties.23

In general, the combination of shape and bond anisotropy leads to an extraordinary control over the crystal structure: polyhedral nanoparticles covered with DNA surface ligands have been shown, for instance, to assemble into crystals fully determined by the size and the symmetry of the particles and by the length of the DNA-ligands. 24,25 DNA-based functionalization can also be achieved by creating suitably shaped frames for spherical nanoparticles: two-dimensional, square-like DNA frames with functionalized edges are able to tune the assembly of the resulting complex units from finite clusters (micelles as well as chain-like aggregates) to planar architectures;<sup>26</sup> while tetravalent DNA-cages are able to induce the assembly of isotropic nanoparticles into diamond crystals.<sup>27</sup>

Within this vast realm, colloidal platelets, i.e. colloids with lateral size at least one order of magnitude smaller than the other two dimensions, may show, e.g., interesting electronic properties as isolated fluorescent emitters28 or even form tilings with photonic properties.<sup>29</sup> In general, colloidal platelets of different shapes can be realized both at the nanometer and the micron scale by, e.g., folding long, single-stranded DNA molecules to create two-dimensional shapes, 30,31 by soft lithography<sup>10</sup> or by making use of preferred growth mechanisms thus obtaining, e.g., silica polygonal truncated pyramids, lanthanide fluoride nanocrystals or polymer-based platelets. 6,14,32,33 Directional bonding can then be added by, e.g., covering the particle edges with ligands<sup>14</sup> or immersing the platelets in a

<sup>&</sup>lt;sup>a</sup> Faculty of Physics, University of Vienna, Boltzmanngasse 5, A-1090, Vienna, Austria. E-mail: carina.karner@univie.ac.at

<sup>&</sup>lt;sup>b</sup> Institut für Theoretische Physik, TU Wien, Wiedner Hauptstraße 8-10, A-1040 Wien, Austria. E-mail: emanuela.bianchi@tuwien.ac.at

<sup>&</sup>lt;sup>c</sup> CNR-ISC, Uos Sapienza, Piazzale A, Moro 2, 00185 Roma, Italy

<sup>†</sup> Electronic supplementary information (ESI) available. See DOI: 10.1039/d0sm00044b

liquid crystal.<sup>6</sup> While the directional bonding constraints favor particle configurations that maximize the number of bonds, the particle shape favors edge-to-edge contacts; the possible competition between these two driving-agents can lead to selfassembled tilings with tunable properties.34-36

Here we consider hard colloidal platelets with a regular rhombic shape decorated with two mutually attractive patches arranged - in different geometries - along two adjacent particle edges. We define two classes of rhombi characterized by having the two patches enclosing either the big or the small angle and, for each of them, we move the patches either in a symmetric or asymmetric fashion along the particle edges. Thus, the resulting building blocks present a wide range of bonding patterns, the combination of which gives rise to different assembly products: chains as well as micelles with three-, fiveor six-fold symmetry. We first investigate the emerging assembly scenarios according to the patch positioning at different interaction energies. For most of the studied systems, we are able to trace a region in the parameter space where micelles are the prevalent assembly product. Hence the second part of our paper investigates if and how systems of micelles with different geometries form a well-defined lattice via hierarchical assembly.

### 2 Methods

#### 2.1 Particle model

We consider regular hard rhombi decorated with two attractive patches. The interaction potential between two of such hard particles, i and j, is given by

$$U(\vec{r}_{ij}, \Omega_i, \Omega_j) = \begin{cases} 0 & \text{if } i \text{ and } j \text{ do not overlap} \\ \\ \infty & \text{if } i \text{ and } j \text{ do overlap} \end{cases}$$

where  $\vec{r}_{ij}$  is the center-to-center vector, while  $\Omega_i$  and  $\Omega_j$  are the particle orientations. Overlaps between rhombi are detected via the separating axis theorem for convex polygons.<sup>37,38</sup>

The patch-patch interaction is a square-well attraction given

$$W(p_{ij}) = \begin{cases} -\varepsilon & \text{if } p_{ij} < 2r_{p} \\ 0 & \text{if } p_{ij} \ge 2r_{p}, \end{cases}$$

where  $p_{ij}$  is the patch-patch distance,  $2r_p$  is the patch diameter and  $\varepsilon$  denotes the patch interaction strength.

A patchy rhombi model of this kind was first detailed in ref. 39, with four attractive patches placed in the center of the edges. In this work, we focus on two-patch rhombi where the patches either enclose the big angle (manta rhombi, referred to as "ma") or the small angle (mouse rhombi, referred to as "mo"); see the particle sketches reported at the top of Fig. 1.

In either type of system, patches can be placed anywhere on the respective edges, resulting in an - in principle - infinite number of possible two-patch rhombi systems. To methodically characterize the patch positioning, we introduce two patch topologies. Patch topologies prescribe how to move the patches with respect to each other. In the symmetric/asymmetric (s/as) topology, patches are placed symmetrically/asymmetrically with

respect to their enclosing vertex. Note that, within a specific topology, the relative distance  $\Delta$  of one patch with respect to the enclosing vertex also determines the position of the other patch: in the s-topology, both patches are placed at distance  $\Delta$  from their common vertex, in the as-topology, one patch is at distance  $\Delta$  from the reference vertex, while the other is at  $(1 - \Delta)$ . With these definitions a two-patch system is fully defined through its patch configuration (ma or mo), its topology (s or as) and its relative position on the edge ( $\Delta$ ). It is important to note that when patches are placed in the edge-center, i.e., at  $\Delta = 0.5$ , the s- and as-topology collapse into the same topology, referred to as center topology, so that the respective rhombi systems are referred to as ma- and mo-center systems. A summary of the used particle parameters can be found in Table 1.

#### 2.2 Simulation details

For the first-stage assembly, we use grand canonical Monte Carlo (GCMC) simulations with standard periodic boundary conditions to model the adsorption of the self-assembling platelets on a surface.<sup>39</sup> Together with single particle rotation/ translation moves and particle insertions/deletions, we implement also cluster moves 40,41 to avoid kinetic traps. We equilibrate the systems for  $3 \times 10^5$  MC-sweeps at a low packing fraction (i.e., at  $\phi \approx 0.05$ ) with  $\mu_{\rm eq}$  and then we increase the chemical potential to  $\mu^*$  to observe the assembly. We run the simulations for about  $\approx 3 \times 10^6$ –7.0  $\times 10^6$  MC-sweeps before collecting statistics. In order to further improve statistics, we perform eight simulations runs per system and interaction strength. For the second-stage assembly - from the aggregated clusters into super-lattices - we run isobaric-isothermal Monte Carlo simulations (NPT-MC) with isotropic volume moves at different pressure values, P. The system parameters for all simulations are given in Table 2.

## 3 Results

In general, the investigated systems yield three different classes of self-assembly products: chains, loops and micelles, emerging from the specific interplay between steric constraints and patchiness, and characterized by specific bonding patterns. By construction, our patchy rhombi can form a maximum of two bonds per particle, one per edge. Each of such bonds can occur either when the rhombi are oriented parallel (p) to each other or when the rhombi are arranged in an arrowhead - i.e., non-parallel (np) - configuration. Chains must have p-bonded but can also present np-bonds, loops can accommodate both p- and np-bonds, while micelles are minimal loops and consist of np-bonds only.

We find that micelles are mostly in competition with chain assemblies, as loops rarely form. While our previous publication detailed the properties of chain assemblies,42 this work focuses on micelles. First, within the framework of a small cluster analysis, we establish for which topologies and patch positions micelles can form (see Fig. 1). Subsequently, we determine for which  $(\Delta, \varepsilon)$ values micelles are the dominant assembly product in simulations

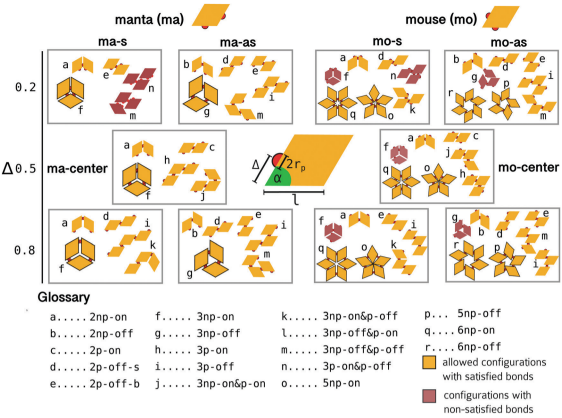


Fig. 1 Sketch of a patchy rhombus (in the center) and small cluster analysis for manta (ma) and mouse (mo) systems in the symmetric (s) and asymmetric (as) topologies. The columns correspond to different particle classes (ma, mo) and different topologies (s, as), as labeled; the rows correspond to different  $\Delta$ -values. At  $\Delta$  = 0.5, the s- and as-topologies collapse to the center topologies, referred to as ma-center and mo-center. Clusters that satisfy bonding restrictions are colored in yellow and clusters that do not satisfy bonding restrictions and would yield non-satisfied bonds are colored in burgundy. Allowed micelles are enlarged and outlined in black. Cluster labels - in the letters a-r - are detailed in the glossary at the bottom. In the glossary, each configuration is characterized by the number of particles in the cluster (from two up to six) and by the bond type: parallel (p) vs. non-parallel (np) as well as on-edge (on) vs. off-edge (off). Thus, dimers with on-edge non-parallel bonds (2np-on) are labeled with a, dimers with off-edge non-parallel bonds (2np-off) are labeled with b, dimers with on-edge parallel bonds (2p-on) are labeled with c, dimers with off-edge parallel bonds (2p-off) are labeled with either d (off-edge bonds closer to the small angle, 2p-off-s) or e (off-edge bonds closer to the big angle, 2p-off-b). The same logic applies to trimers and bigger clusters.

**Table 1** Particle parameters: the opening angle  $\alpha$ , the side length l (which sets the unit of length), the patch size  $2r_p$ , the patch-patch attraction strength  $\varepsilon$  (in unit of  $k_BT$ ), the patch position  $\Delta$ , as labeled. Note that  $\Delta$ refers to the relative placement of a patch on a rhombi edge. See Fig. 1 for sketches

Parameter	Symbol	Value
Angle Side length Patch radius Interaction strength Patch position	$egin{array}{c} lpha \ l \ r_{ m p} \ arepsilon \ arepsilon \ arDelta \ \end{array}$	60° 1.0 0.05 [5.2, 10.2] [0.2, 0.8]

(see Fig. 2-4). Finally, we compress the micelle assemblies with the highest yield and analyze the quality of the second-stage assembly products (see Fig. 5-7).

#### 3.1 Small cluster analysis

The small cluster analysis reported in Fig. 1 allows us to discern which clusters fulfill the given bonding constraints irrespective of their formation probability in a simulation or experiment.

Table 2 System parameters in reduced units (as used in all simulations)

System parameter	Symbol	Value
Area of simulation box	A	1000·sin(60°)
Box width	$L_x$	$\sqrt{1000}$
Box height	$L_{y}$	$\sqrt{1000} \cdot \sin(60^\circ)$
Chemical potential eq.	$\mu_{ m eq}$	0.1
Chemical potential	$\mu^{\star}$	0.3
Boltzmann constant	$k_{ m B}$	1
Temperature	T	0.1
Pressure	P	[5, 100]

Note that the bonding constraints themselves follow from the patch configuration (ma or mo), the patch topology (s or as) and the patch position ( $\Delta$ ). In addition to p- and np-bonds, we distinguish between on-edge bonds (on) where the edges align and off-edge bonds (off) where the edges are offset with respect to each other. In the case of the p-off bonds, we further specify if the bond is closer to the small (p-off-s) or to the big angle (p-off-b) (see the small cluster analysis and the glossary of Fig. 1 for sketches of bonding scenarios).

**Paper** Soft Matter

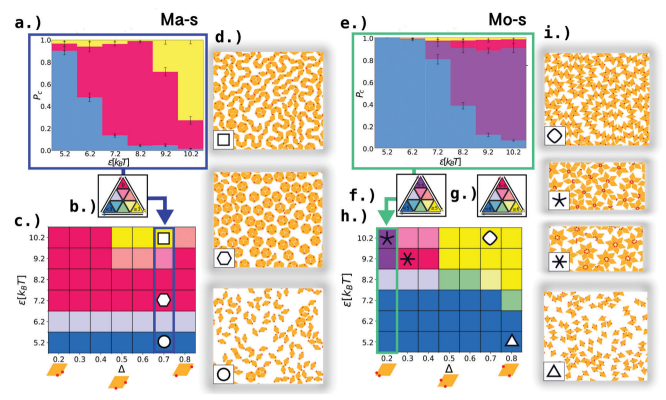


Fig. 2 Self-assembly products of symmetric manta (ma-s) and mouse (mo-s) systems. (a) Histogram of yields of different cluster types in the ma-s system at  $\Delta = 0.7$  and  $\varepsilon = [5.2, 10.2]k_BT$ . Histograms for the other  $\Delta$ -values can be found in Fig. S1 of the ESI.† Cluster types of interest are: clusters with size N < 3 (liquid, blue), three np-bonded particle loops (boxes, pink) and clusters with  $N \ge 3$  (chains/loops, yellow). The barycentric color triangle in (b) maps the distribution of cluster types for all  $(\Delta, \epsilon)$ -values to the heatmap in (c) (see the ESI† for the detailed description of the barycentric coordinates). (d) Simulation snapshots of ma-s. From top to bottom: chains, loops and boxes at  $\varepsilon = 10.2k_BT$ ,  $\Delta = 0.7$ , boxes at  $\varepsilon = 7.2k_BT$ ,  $\Delta = 0.7$ , rhombi liquid at  $\varepsilon = 5.2k_BT$ ,  $\Delta = 0.7$ . (e) Histogram of yields of different cluster types in the mo-s system at  $\Delta = 0.2$  and  $\varepsilon = [5.2, 10.2]k_BT$ . Histograms for the other  $\Delta$ -values can be found in Fig. S2 of the ESI.† Cluster types of interest are: clusters with N < 5 (liquid, blue), five np-bonded particle loops (5-stars, purple), 6 np-bonded particle loops (6-stars, pink) and clusters with  $N \ge 5$  (chains/loops, yellow). The barycentric color triangles in (f) and (g) map the distribution of cluster types to the heatmap column/s for  $\Delta = 0.2/[0.3, 0.8]$  in (h). (i) Simulation snapshots of mo-s. From top to bottom: chains, loops and stars at  $\varepsilon=10.2k_{B}T$ ,  $\Delta=0.7$ , 5-stars at  $\varepsilon=10.2k_{B}T$ ,  $\Delta=0.2$ , 6-stars at  $\varepsilon=9.2k_{B}T$ ,  $\Delta=0.3$ , rhombi liquid at  $\varepsilon=5.2k_{B}T$ ,  $\Delta=0.8$ .

In ma-s, micelles consist of three np-on-bonded particles (also referred to as boxes) and they can form at all  $\Delta$ -values (see clusters labeled as f in the corresponding panels of Fig. 1). For  $\Delta$  < 0.5 micelles are the only possible self-assembly product as bonding incompatibilities prevent dimers (see cluster e in the corresponding panel of Fig. 1) from growing into chains and loops (see clusters n and m, respectively). In contrast, for  $\Delta \geq 0.5$ , chains and loops can form since the patch positioning does not anymore disfavor the formation of p- and mixed-bonded clusters of sizes bigger than two (see clusters h, i, j and k in the corresponding panels of Fig. 1).

In mo-s, micelles consist either of five or six np-on-bonded particles (referred to as 5- and 6-stars, respectively) and they can form at all ∆-values (see clusters o and q in the corresponding panels of Fig. 1). We note that, micelles of three particles (boxes) would have non-satisfied bonds, thus they are energetically disfavored with respect to the stars. For  $\Delta$  < 0.5, threeparticle clusters with p- and np-bonds are still allowed (see cluster k in the corresponding panel of Fig. 1), thus rendering chains a possible self-assembly product, albeit with a restricted configuration space. For  $\Delta \geq 0.5$ , also both p- and mixed-bonded

clusters can form (see clusters h, i, j and k in the corresponding panel of Fig. 1).

In both ma-as and mo-as, there are no bonding restrictions for p-bonds, and hence chains, loops and micelles are allowed for all values of  $\Delta$ . Furthermore, micelles have pores in the center as particles can form np-bonds only off-edge. Therefore, the resulting clusters are referred to as open boxes in the case of ma-as (see clusters g in the corresponding panels of Fig. 1) and open stars in the case of mo-as (see clusters p and r in the corresponding panels of Fig. 1).

It is important to note that while in s-topologies the geometrical form of the assembled micelles is the same at all △-values, in as-topologies the cluster geometry does change because of the  $\Delta$ -dependent opening and closing of the pores. While at  $\Delta = 0.5$  (corresponding to ma- and mo-center), the pores are completely closed yielding closed boxes and stars, the pores continuously open as  $\Delta$  is moved off-center and – as a result – the rhombi corners protrude more and more. As in ref. 34, in the sticky limit, the side length of the pores is given by

$$l_{\text{pore}} = |l - 2\Delta|. \tag{1}$$



Ma-as Mo-as a.) e.) i.) d.) 1.0 0.8 0.6 0 10.2 7.2 8.2 ε[k<sub>B</sub>T] 9.2  $\varepsilon[k_BT]$ \* c.) **h.**)<sub>10.2</sub> 10.2 9.2 X 8.2 \* 8.2 7.2 7.2 6.2 6.2 5.2

Fig. 3 Self-assembly products of asymmetric manta (ma-as) and mouse (mo-as) systems. (a) Histogram of yields of different cluster types in the ma-as system at  $\Delta = 0.7$  and  $\varepsilon = [5.2, 10.2]k_BT$ . Histograms for remaining  $\Delta$  values can be found in Fig. S3 of the ESI.† Cluster types of interest are: clusters with size N < 3 (liquid, blue), three np-bonded particle loops (open boxes, pink) and clusters with  $N \ge 3$  (chains/loops, yellow). The barycentric color triangle in (b) maps the distribution of cluster types for all  $(\Delta, \varepsilon)$ -values to the heatmap in (c) (see the ESI† for the detailed description of the barycentric coordinates). (d) Simulation snapshots of ma-as. From top to bottom: chains, loops and open boxes at  $\varepsilon = 10.2k_{\rm B}T$ ,  $\Delta = 0.7$ , open boxes at  $\varepsilon = 8.2k_{\rm B}T$ ,  $\Delta = 0.7$ , rhombi liquid at  $\varepsilon = 5.2k_BT$ ,  $\Delta = 0.7$ . (e) Histogram of yields of different cluster types in the mo-as system at  $\Delta = 0.2$  and  $\varepsilon = [5.2, 10.2]k_BT$ . Histograms for the other  $\Delta$ -values can be found in Fig. S4 of the ESI.† Cluster types of interest for  $\Delta$  = 0.2 are: clusters with size N < 5 (liquid, blue), five np-bonded particle loops (open 5-stars, purple), six np-bonded particle loops (open 6-stars, pink), and clusters with  $N \ge 5$  (chains/loops, yellow). The barycentric color triangle in (f) and (g) maps the distribution of cluster types to the heatmap column/s for  $\Delta = 0.2/[0.3, 0.8]$  in (h). (i) Simulation snapshots of mo-as. From top to bottom: chains, loops and open stars at  $\varepsilon = 10.2k_BT$ ,  $\Delta = 0.2$ , chains, loops and open stars at  $\varepsilon = 8.2k_BT$ ,  $\Delta = 0.2$ , chains, loops and open stars at  $\varepsilon = 8.2k_BT$ ,  $\Delta = 0.3$ , rhombi liquid at  $\varepsilon = 5.2k_BT$ ,  $\Delta = 0.2$ .

0.3 0.4

0.5 0.6

Δ

0.7 0.8

Once  $l_{\text{pore}}$  is known, the area of the pores can be calculated for both triangular (ma-as systems) and hexagonal pores (mo-as systems).34

0.7 0.8

#### 3.2 First stage self assembly

0.2 0.3 0.4 0.5 0.6

Beyond the described bonding restrictions, we need to determine under which conditions the different aggregates emerge in many-body systems. For that, we run GCMC simulations and, at the end of the self-assembly process, we calculate yields of chains/loops and micelles.

The yield of a cluster type is defined as the percentage of particles in clusters belonging to the selected cluster type. We define cluster types of interest according to the specific systems. In ma-systems - for both the s- and the as-topology we distinguish clusters of size N < 3 (classified as liquid), three np-bonded particle loops (classified as boxes/open boxes) and non-box clusters with size  $N \ge 3$  (corresponding to chains/ loops). In mo-systems – both s and as – we distinguish clusters with size N < 5 (liquid), five np-bonded particle loops (5-stars/ open 5-stars), six np-bonded particle loops (6-stars/open 6-stars) and non-star clusters  $N \ge 5$  (chains/loops).

The obtained yields are reported as a function of  $\varepsilon$  for each △-value (see panels a and e of Fig. 2 and 3 for an example  $\Delta$ -value, while for the full overview of  $\Delta$ -values see Fig. S1–S4 in the ESI†). Through a mapping to a barycentric coordinate system (see ESI† for the mapping and calculation details), we obtain heatmaps for the four investigated systems (ma-s, mo-s, ma-as and mo-as) where at each  $(\Delta, \varepsilon)$ -value we identify the predominant self-assembly scenario. The heatmaps reflect the relative dominance of the different cluster types: whenever one cluster type has a yield higher than 2/3, the heatmap obtains the color of the respective triangle edge (blue for liquid, yellow for chains/loops, pink and purple for micelles) and we call this cluster type 2/3-dominant; if no cluster type dominates by more than 2/3, one of the mixed colors in the center of the triangle is adopted (see panels c and h of Fig. 2 and 3 for the heatmaps, see panels b, f and g in Fig. 2 and 3 for the respective barycentric triangles). Simulation snapshots are displayed in Fig. 2d and i for ma-s/mo-s systems and in Fig. 3d and i for ma-as/mo-as systems. Micelle yields for all systems are given in Fig. 4.

Paper

a.)ma-s d.)ma-as 0.8 0.8 0.6 0.6 yield yield open boxes boxes 0.2 0.2 0.001 0.00.2 mo-as b.)mo-s 0.8 0.8 0.6 0.6 5-stars <u>0</u>0.4 open 5-stars 0.2 0.2 0.00.2 0.8 0.8 c.) mo-s mo-as 0.8 0.8 6-stars 0.6 0.6 open 6-stars <u>9</u>0.4

Fig. 4 Yields of micelles as a function of  $\Delta$ : (a) boxes in ma-s systems. (b) 5-Stars in mo-s systems. (c) 6-Stars in mo-s systems. (d) Open boxes in ma-as systems. (e) Open 5-stars in mo-as systems. (f) Open 6-stars in mo-as systems. Different curves within each panel denote yields at different  $\varepsilon$ -values and are colored according to the legend at the bottom. Yields are averaged over eight simulation runs per system.

 $\varepsilon =$ 

 $5.2k_BT$ 

0.2

 $7.2k_BT$ 

 $8.2k_BT$ 

 $9.2k_BT$ 

 $10.2k_BT$ 

**ε** =

We start our discussion with ma-s systems (Fig. 2, panels from a to d). When  $\varepsilon \leq 6.2k_{\rm B}T$  such systems remain liquid over the whole  $\Delta$ -range. On increasing  $\Delta$ , the self-assembly process takes place: at  $\varepsilon = 7.2k_{\rm B}T$  and  $\varepsilon = 8.2k_{\rm B}T$ , boxes are the 2/3-dominant self-assembly product at all ∆-values; while for  $\varepsilon \geq 9.2k_{\rm B}T$  there exists a range of  $\Delta$ -values where chains/loops prevail over the micelles (see Fig. 2c). The yield of boxes increases monotonically with  $\varepsilon$  for all  $\Delta$ -values until  $\varepsilon = 9.2k_{\rm B}T$ (see Fig. 4a). For  $\varepsilon \geq 9.2k_{\rm B}T$  and  $\Delta < 0.5$ , the yield of the boxes reaches values above 0.95, while at  $\Delta = 0.5$ , we observe a sudden drop: for  $\Delta \geq 0.5$  bonding restrictions allow chains/loops to form and compete with boxes and as soon as the interaction strength is  $\varepsilon \geq 9.2k_{\rm B}T$ , chains/loops prevail over the boxes. The dip in the box-yield at ma-center becomes deeper on increasing  $\varepsilon$  and while for  $\varepsilon = 9.2k_{\rm B}T$  the yield recovers to 0.8 at  $\Delta = 0.8$ , for  $\varepsilon = 10.2k_{\rm B}T$  the yield remains below 0.45 at  $\Delta = 0.8$ . Summarizing, in ma-s, boxes prevail over chains/loops over a wide range of  $(\Delta,\varepsilon)$ -values.

In mo-s systems (Fig. 2, panels from e to i), we observe characteristic self-assembly products only for  $\varepsilon \geq 8.2k_{\rm B}T$ . At such high interaction strengths, micelles emerge with a non-negligible yield at  $\Delta=0.2$  in the case of 5-stars and at  $\Delta=0.3$  and 0.4 in the case of 6-stars. For these  $\Delta$ -values, the staryields rise monotonically with  $\varepsilon$  (see Fig. 4b for the 5-star-yield and

Fig. 4c for the 6-star-yield), but only reach values above 2/3 for  $\varepsilon \ge 9.2k_{\rm B}T$  (see Fig. 2h). The highest 5-star-yield is reached at  $\varepsilon = 10.2k_{\rm B}T$  and  $\Delta = 0.2$  with  $0.838 \pm 0.04$ . On increasing  $\Delta$ , the 5-star-yield falls to  $0.084 \pm 0.018$  at  $\Delta = 0.3$  and becomes smaller than 0.025 at  $\Delta = 0.5$ . In contrast, 6-stars reach their highest yields at  $\varepsilon = 9.2k_{\rm B}T$ , with the highest value  $0.754 \pm 0.028$  at  $\Delta = 0.4$ . On increasing  $\Delta$ , the 6-star-yield drops to  $\approx 0.2$  at mo-center and proceeds to drop gradually below 0.05 at  $\Delta = 0.8$ . With star-yields below 0.2 for  $\Delta \ge 0.5$ , chains/loops prevail as dominant clusters at high  $\varepsilon$ -values (see Fig. 2h for phase boundaries).

Summarizing, in mo-s, stars have a significantly narrower  $(\Delta, \varepsilon)$ -range of prevalence compared to boxes in ma-s (see Fig. 2c for ma-s and Fig. 2h for mo-s). While boxes are 2/3-dominant for all  $\Delta$  at intermediate  $\varepsilon$ -values, stars reach yields above 2/3 only at high  $\varepsilon$ -values and only for  $\Delta$  < 0.5. At high  $\varepsilon$  and for  $\Delta$  < 0.5 extended chains become available to mo-s and are formed at a higher rate.

In ma-as systems (Fig. 3, panels from a to d), the yield of micelles (open boxes, in this case) rises monotonously with  $\varepsilon$ : open boxes become 2/3-prevalent for intermediate interaction strengths (*i.e.*, at  $\varepsilon = 7.2k_{\rm B}T$  and  $8.2k_{\rm B}T$ ) over the whole  $\Delta$ -range, with the highest yields – above 0.9 – achieved at  $\varepsilon = 8.2k_{\rm B}T$  (see Fig. 3c for the heatmap and Fig. 4d for numerical values of yields). At high interaction strengths, *i.e.*, for  $\varepsilon \geq 9.2k_{\rm B}T$ , the open box yield drops for all  $\Delta$ , first to 0.4 at  $\varepsilon = 9.2k_{\rm B}T$  and then to 0.2 at  $\varepsilon = 10.2k_{\rm B}T$ . The drop in the micelle-yield is due to the emergence of chains, that can form at all  $\Delta$ -values and prevail over the open boxes when  $\varepsilon \geq 9.2k_{\rm B}T$ .

In mo-as systems (Fig. 3, panels from e to h), micelles (open stars, in this case) do not become dominant at any  $(\Delta, \varepsilon)$ -value. In particular, the yield of open 5-stars stays below 0.03 for all  $\Delta$ -values and does not increase with  $\varepsilon$ , meaning that these micelles are always negligible compared to chains/loops (see Fig. 4e). In contrast, the yield of open 6-stars does increase with  $\varepsilon$  for all but the most extreme  $\Delta$ -values (i.e.,  $\Delta$  = 0.2 and 0.8); nonetheless, as the maximum yield stays below 0.25 (see Fig. 4f), also these micelles are negligible over the whole  $\Delta$ -range. While for  $\varepsilon \leq 7.2k_{\rm B}T$ , the systems are mostly liquid, for  $\varepsilon \geq 7.2k_{\rm B}T$ , chains become the most dominant cluster type at all  $\Delta$ -values.

Concluding, micelles in as-topologies have a smaller window of prevalence compared to micelles in s-topologies because in as-topologies extended p- and mixed bonded chains are allowed at all  $\Delta$ -values and they dominate over micelles for high interaction strengths. In the following, we describe the properties of the observed micelles, mostly focusing on their packing properties. For the detailed characterization of the chain/loops assemblies we refer the reader to ref. 42.

#### 3.3 Second-stage assembly

To further explore the versatility of patchy rhombi as building blocks, we compress configurations with high micelle yields and thereby induce a second-stage assembly process. We use the configurations as they formed within simulations, including all not fully formed and misshaped clusters and monomers. This enables us to study the efficiency of the hierarchical assembly

ma-s<sub>Δ=0.3</sub> b.)  $\Delta = 0.7$ 100  $\Delta = 0.5$  $\Delta = 0.3$ 0.3 80  $P=10, \Delta=0.3, S_1=0.032$ 60 0.2 40 0.1  $P=30, \Delta=0.3, S_1=0.130$ 20 0 0.0 0.6 0.7 0.4 0.5 0.8  $P=100, \Delta=0.3, S_1=0.373$ c.) d.) boxes,  $\Delta=0.3,\,P=100$ 135 45  $\langle S_L^2 \rangle$ 180

Fig. 5 Second-stage assembly of boxes at  $\epsilon=8.2k_{\rm B}T$  for symmetric manta systems (ma-s). (a) Left: Simulation snapshots of the system with  $\Delta=0.3$  at P=[10,30,100] (from top to bottom). Right: Centers of mass of boxes of the respective snapshots colored according to the hexatic order parameter,  $\Psi$ , defined in eqn (3) (see the color wheel in panel c). (b) Eos for box systems at  $\Delta=[0.3,0.5,0.7]$ , where  $\phi$  denotes the packing fraction defined in eqn (2). (c) Color wheel denoting the values of  $\Psi$  for the center of mass snapshots in panel a. (d) Average fraction of the largest hexagonal domain  $\langle S_L \rangle$  — defined in the text — as function of P; note that the average is taken over all  $\Delta$ -values; note that the background of the eos-plot in panel a is colored according to  $\langle S_L \rangle$ . Error bars denote the standard deviation and their wide extent is due to relatively small system sizes. (e) The radial distribution function g(r) at  $\Delta=0.3$  and P=100.

P

100

process, from two-patch rhombi to micelle lattices and show viable routes for material synthesis. Within our computer simulations, the second-stage self-assembly is modeled with *NPT*-MC simulations and the pressure range is P=5–100. We selected systems with a high percentage of micelles, *i.e.*, with yields higher than 0.75. While there are  $(\Delta, \varepsilon)$ -values for which boxes, open boxes and stars reach yields above 0.75, maximum yields of open stars stay below 0.25. Therefore we exclude them from our second-stage assembly investigation and focus on boxes, open boxes and stars. Note that boxes, open boxes and stars are hard particles: boxes are convex as they are regular hexagons; open boxes are non-convex, with their protrusions growing more prominent as  $\Delta$  moves towards off-center-values; 5-stars and 6-stars are non-convex.

270°

Upon increasing pressure, micelles pack and tend to form crystalline domains with local hexagonal order. We define the packing fraction as

$$\varphi = N_{\rm p} \frac{A_{\rm p}}{4},\tag{2}$$

0.2

where  $N_{\rm p}$  is the total number of particles,  $A_{\rm p}$  is the area of a rhombic platelet and A is the total area of the simulation box. We note that the area of the patches is not included in  $A_{\rm p}$  as the patches can completely overlap. The equations of state (eos) for the different systems at selected  $\Delta$ -values are reported in panel b of Fig. 5 (ma-s), Fig. 6 (ma-as) and Fig. 7 (mo-s). To characterize the emergence of long range order, we calculate the radial distribution function, g(r), where r is the center-to-center distance between two micelles. At this point we note that the side length of our rhombi is l=1. Additionally, we measure the local order with the hexatic order parameter, that is given for every micelle j as

$$\Psi_j = \frac{1}{N_n} \sum_{k=1}^{N_n} \exp(i6\Theta_{j,k}), \tag{3}$$

where  $N_n$  is the number of all next neighbors and  $\Theta_{j,k}$  is the relative bond angle between j and its neighbor micelle k, with respective to a chosen simulation box vector. To determine all

**Paper** 

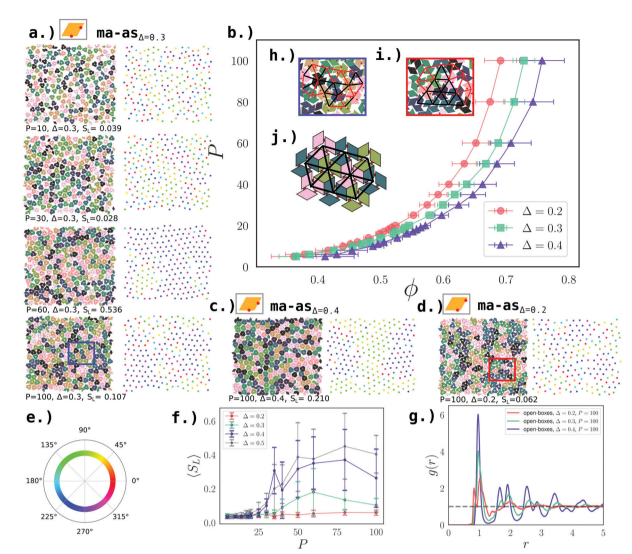


Fig. 6 Second-stage assembly of open boxes at  $\varepsilon = 8.2k_BT$  for asymmetric manta systems (ma-as). (a) Left: Simulation snapshots of the system with  $\Delta = 0.3$  at P = [10, 30, 60, 100] (from top to bottom); clusters are colored to facilitate visual distinction. Right: Centers of mass of open boxes of the respective snapshots colored according to the hexatic order parameter,  $\Psi$ , defined in eqn (3) (see the color wheel in panel e). (b) Eos for open box systems at  $\Delta = [0.2, 0.3, 0.4]$ , where  $\phi$  is the packing fraction defined in eqn (2). (c) Left: simulation snapshot of the system at  $\Delta = 0.4$  and P = 100. Right: Centers of mass of open boxes of the respective snapshot colored according to  $\Psi$ . (d) Left: Simulation snapshot of the system at  $\Delta = 0.2$  and P = 100. Right: Centers of mass of open boxes of the respective snapshot colored according to  $\Psi$ . (e) Color wheel for the values of  $\Psi$  for the center of mass snapshots in panels a, c and d. (f) Average fraction of largest hexagonal domain  $\langle S_L \rangle$  – defined in the text – as function of P for  $\Delta$  = [0.2, 0.3, 0.4, 0.5], as labeled. The error bars denote the standard deviation and are wide due to relatively small system sizes. (g) The radial distribution function g(r) at  $\Delta = [0.2, 0.2]$ 0.3, 0.4] and P = 100, as labeled. (h) Magnified view of the blue rectangle of snapshot of ma-as<sub>A</sub> = 0.3, P = 100. Black lines connect boxes at hexagonal lattice positions; red lines connect boxes that are shifted due to defects. (i) Magnified view of the red rectangle of ma-as  $_{1}$  = 0.2. Black lines connect boxes at hexagonal lattice positions; red lines connect boxes that are shifted due to defects. (j) Sketch of the perfect hexagonal open box tiling with triangular pores (ot-tiling) as observed in local regions in simulation snapshots.

next neighbours, we calculate a distance histogram including the first 12 neighbours for every micelle and we define next neighbours as all micelles within the first minimum of this histogram. Finally, to quantify the transition between isotropic to hexatic, we calculate the average fraction of particles in the largest hexagonal domain  $\langle S_L \rangle$ . For the calculation of  $\langle S_L \rangle$ , we consider two neighbouring micelles, i and j as part of the same domain if the difference of their hexatic order parameter  $d\Psi_{ij} < 15^{\circ}$ . The average is then taken over parallel simulation runs.

Box assembly. To study the second-stage assembly of boxes, we select ma-s systems with  $\Delta = [0.3, 0.5, 0.7]$  and  $\varepsilon = 8.2k_BT$ . The box-yields for these systems are 0.963  $\pm$  0.010 at  $\Delta$  = 0.3,  $0.948 \pm 0.011$  at  $\Delta = 0.5$  and  $0.947 \pm 0.016$  at  $\Delta = 0.7$ .

Upon increasing pressure, boxes assemble into a lattice with hexatic/hexagonal order at all selected \( \Delta\)-values. On the left hand side of Fig. 5a, we display simulation snapshots for boxes with  $\Delta = 0.3$  at different pressures, while on the right hand side we show the same snapshots with the box centers colored according to their respective  $\Psi$  (see Fig. 5c for  $\Psi$ -color wheel).

**mo-s**<sub>Δ=0.2</sub>  $mo-s_{\Delta=0.3}$ b.) 100 h.) g.) 80  $P=10, \Delta=0.2$  $P=10, \Delta=0.3$ 60 40  $\Delta = 0.2$ 20  $\Delta = 0.3$  $\Delta = 0.4$  $P=25, \Delta=0.2$  $P=25. \Delta=0.3$ 0 0.5 0.6 0.7 Φ e.)  $P=100. \Delta=0.2$  $P=100, \Delta=0.3$ 0.8 f.) d.) stars,  $\Delta = 0.2$  , P = 100stars.  $\Delta = 0.3$  . P = 100yield [%] 0.4 4 6-star,  $\Delta = 0.2$ 3 5-star,  $\Delta = 0.3$ 0.2 0.0 0 n 50 100 6 6

Fig. 7 Second-stage assembly of 5-stars and 6-stars at  $\varepsilon=9.2k_{\rm B}T$  for symmetric mouse systems (mo-s). (a) Simulation snapshots of the system at  $\Delta=0.2$  at P=[10, 25, 100] (from top to bottom), where 5-stars are predominant. (b) Eos for star systems at  $\Delta=[0.2, 0.3, 0.4]$ , where  $\phi$  is the packing fraction defined in eqn (2). (c) Simulation snapshots of the system at  $\Delta=0.3$  at P=[10, 25, 100] (from top to bottom), where 6-stars are predominant. (d) The radial distribution function g(r) for the 5-star system at  $\Delta=0.2$  and P=100. (e) The yield of 5-stars and 6-stars at  $\Delta=0.2$  and  $\Delta=0.3$  as function of P, as labeled. (f) The radial distribution function g(r) for the 6-star system at  $\Delta=0.3$  and  $\Delta=0$ 

P

We note that, while at P = 10, the system of boxes is liquid, at  $P \approx 20$  the system becomes hexatic, *i.e.*, with local hexagonal order but no long range order. As P increases further, long range order becomes stronger. At P = 100, remaining single rhombi induce grain boundaries into the otherwise perfectly hexagonally ordered domains. We confirm the hexatic/hexagonal order at P = 100 with the radial distribution function (see Fig. 5e). Further, we calculate  $\langle S_{\rm L} \rangle$  as function of the pressure (see Fig. 5d) to visualize the growth of ordered domains we colored the background of the eos-curves (see Fig. 5b). The same scenario is observed at all selected  $\Delta$ -values: from  $\Delta$  = 0.3 to  $\Delta$  = 0.7 the equations of state collapse onto one curve (see Fig. 5b). The maximum packing at the highest pressure P = 100 is  $\phi$  = 0.793  $\pm$  0.006 for all  $\Delta$ -values. Therefore we conclude that for these boxes, the specific ∆-value does not influence the second-stage assembly.

It is important to note that although significant parts of the systems are ordered hexagonally at high pressures, the small system sizes render it impossible to determine whether the system is hexagonal on a long range scale. See ref. 43 for detailed studies on the liquid-hexatic-hexagonal phase transitions.

Open box assembly. For the assembly of open boxes into super structures, we choose ma-as systems at  $\varepsilon=8.2k_{\rm B}T$  and  $\varDelta=[0.2,~0.3,~0.4]$ . Note that at  $\varDelta=0.5$  ma-as collapses to macenter, thus forming boxes (already discussed in Fig. 5). As  $\varDelta$  moves towards off-center values, pores in the boxes open up and become bigger and bigger, while at the same time the corners protrude more and more. As a consequence, the eos-curves gradually shift to lower packing fractions at intermediate and high P-values (see Fig. 6b). While boxes at  $\varDelta=0.5$  and P=100 yield a packing of 0.793  $\pm$  0.006, for open boxes at  $\varDelta=0.3$  the packing is 0.728  $\pm$  0.017, and at  $\varDelta=0.2$  it is 0.691  $\pm$  0.025.

As with the boxes, we also measure the positional order with g(r),  $\langle S_{\rm L} \rangle$  and  $\Psi$  for the open boxes. The peak pattern of the g(r)shows that hexagonal order is still pronounced at  $\Delta = 0.4$ , significantly deteriorates already at  $\Delta = 0.3$  and ceases completely at  $\Delta = 0.2$  (see Fig. 6g). Similarly,  $\langle S_{\rm L} \rangle$  indicates smaller ordered domains for  $\Delta = 0.3$  and  $\Delta = 0.2$  (note that due to large error-bars - that are a result of the relatively small system sizes – we can only report trends for  $\langle S_1 \rangle$ ). The progressive reduction of the hexatic/hexagonal order with increasing  $\Delta$  can also be directly observed in the simulation snapshots reported in Fig. 6a-d as well as in the insets 6h and i. In conclusion, in ma-as the hexatic/hexagonal order gets destroyed as  $\Delta$  shifts towards off-center values.

Star assembly. For the second-stage assembly of stars, we select systems at  $\varepsilon = 9.2k_{\rm B}T$  with  $\Delta = 0.2$  (5-stars) and  $\Delta = [0.3,$ 0.4] (6-stars). The star yields of these initial configurations are  $0.838 \pm 0.041$  for 5-stars at  $\Delta = 0.2, 0.727 \pm 0.025$  for 6-stars at  $\Delta$  = 0.3 and 0.754  $\pm$  0.028 for 6-stars  $\Delta$  = 0.4. It is important to note that these yields from CGMC simulations decay for all NPT-MC runs, even at low pressures P (see Fig. 7e).

The eos-curve for 5-stars (red dots in Fig. 7b) lies slightly above the 6-star curves for all intermediate and high P-values, indicating that 5-stars pack less close at the same P with respect to 6-stars (green squares and blue triangles in Fig. 7b). The packing fraction for 5-stars at P = 100 is 0.696  $\pm$  0.0214, while for 6-stars at  $\Delta = 0.3$  it is  $0.706 \pm 0.018$ .

The analysis of the g(r) at P = 100 (panels d and f of Fig. 7) suggests that in both star systems there exists a residual hexagonal order. However, the resulting structures can not be described as hexagonal lattices due to the many non-star clusters present already in the initial configuration - as the star-yields are low compared to those observed for the boxes. Simulation snapshots in Fig. 7a and c show that these clusters often have the structure of non-completed or misshaped stars. These "broken stars" induce defects to an extent that destroys local and, subsequently, long range order.

Moreover, the 5-star-yield reduces on increasing pressure (see Fig. 7e): while the yield of 6-stars at  $\Delta = 0.3$  only slightly decreases with P, the 5-star yield at  $\Delta = 0.2$  decreases from  $0.75 \pm 0.03$  at P = 10 to  $0.54 \pm 0.04$  at P = 100. This is due to the fact that, as the pressure is increased, the void where a sixth star would fit, starts to fill up. The result are conglomerates of destroyed 5-stars (see Fig. 7b, red inset). Together with already present grain boundaries, these conglomerates completely break the hexatic/hexagonal lattice order.

In contrast, for 6-stars, we observe the presence of residual areas with hexagonal ordering and triangular pores independent of  $\Delta$  (see Fig. 7b, dark blue inset).

#### 4. Conclusion

In this paper, we investigate the assembly of patchy colloidal platelets with a regular rhombic shape. We consider a simple, purely two-dimensional model consisting of rhombi decorated with two bonding patches, arranged in different ways along

adjacent particle edges. Rhombic platelets realized, for instance, with shape anisotropic DNA origami could be decorated with sticky-ended DNA strands: the possibility to place them with high accuracy on the platelets surface<sup>27,30</sup> would allow the reproduction of different patch topologies.

We explore dynamically accessible self-assembly products at finite temperatures to understand which self-assembly scenarios would emerge in experiments. Depending on the patch arrangement and interaction strength we find micelles (minimal loops) or chains to be the prevalent self-assembly product. We observe that equivalent simulation runs produce comparable chain and micelle yields over simulation times of about 10<sup>7</sup> MC sweeps.

While an earlier work studied the properties of chains, 42 this work focuses on micelles. In general, we distinguish two types of micelles: boxes, that emerge for systems where the patches enclose the big angle (manta), and stars that are formed by systems where the patches enclose the small angle (mouse).

Boxes and stars can be stabilized for systems where both patches are distributed symmetrically at a distance less than 0.5  $(\Delta < 0.5)$  from their enclosing angle. For systems with  $\Delta \geq 0.5$ , micelles and chains compete at high interaction strengths.

In systems, where patches are distributed asymmetrically *i.e.*, the distance of one patch to the enclosing angle is  $\Delta$ , while the distance of the second patch is  $(1 - \Delta)$  - stars and boxes have a pore in the center and we call them open boxes and open stars. We note that while for large as well as small \( \Delta\)-values the pore size is maximal, the pore closes off completely for  $\Delta = 0.5$ and symmetric and asymmetric patch distributions are identical. We find that, while open boxes are prevalent at intermediate interaction strength, open stars never reach a yield over 0.25. At high interaction strength chains prevail for all △-values in both, manta and mouse systems.

We note that in our investigations the patch size is fixed to 1/10th of the side length. As this feature can be tuned in experimental systems, for instance by changing the length of the stickyended DNA strands, it would be interesting to investigate how the self-assembly process is influenced by a change in the patch size. From our results we speculate that, a larger yield of chains would be observed on increasing the patch size, as the larger patch areas would imply a stronger interaction strength.

In the second part of the paper we compress the systems of boxes, open boxes and stars and characterize the emerging lattices. Of course, the yield of the first structures plays an important role in the second-stage assembly process, 19 so we compress only those systems where micelle yield is sufficiently high.

While boxes yield a hexagonal lattice independent of  $\Delta$ , open boxes self-assemble into a hexagonal lattice with a  $\Delta$ -dependent long range order. As  $\Delta$  becomes more extreme, gaps and pores within the lattice grow larger, while the long range order decays. In contrast, the self-assembly product of stars displays no long range hexagonal order for any ∆-value.

We note that, in box systems defects in the hexagonal order are exclusively due to either vacancies or not fully formed boxes. In open box and star systems, on the other hand, defects inherent to the particle shape arise.

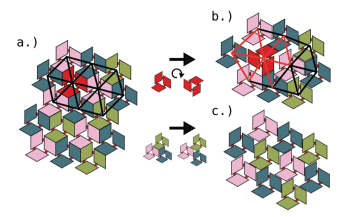


Fig. 8 Tilings of open boxes where the clusters are depicted in different colors to facilitate visual distinction. (a) Perfect hexagonal tiling with triangular pores, referred to as open triangular tiling (ot-tiling). The black lines connect the centers of neighboring boxes and highlight the hexagonal order of the tiling. (b) Representation of a lattice defect: the red open box is rotated by 60° with respect to the perfect ot-tiling in panel a. Due to the rotation, the rhombi of the red open box are aligned parallel with their neighboring rhombi belonging to a different box. The hexagonal lattice is thus distorted: while neighboring boxes in correct lattice positions are still connected by black lines, boxes that had to shift due to the orientational defect are connected with red lines. (c) Perfect hexagonal tiling of open boxes with triangular and hexagonal pores, referred to as open triangular-hexagonal tiling (oth-tiling). This tiling - also known as kagome lattice - was found in ref. 34 in systems of patchy rhombi with four patches.

For what concerns open boxes, the best hexagonal order would be achieved by the lattice reported in Fig. 8a. We refer to this lattice as open triangular tiling (ot-tiling) as it has triangular pores, whose size depend on  $\Delta$ . In the ot-tiling, all rhombi are aligned in a non-parallel fashion. Although such an alignment is able to incorporate the protruding corners while maximizing edge-to-edge contact, other on-edge alignments are possible and can poison the local order. As a consequence, in simulations, we observe orientational point defects due to the fact that rhombi belonging to different boxes align in a parallel way, as shown in Fig. 8b. While for closed boxes this kind of parallel alignment is commensurable with the closed-packed hexagonal lattice, in open box systems, lattice distorting defects are introduced and the more extreme  $\Delta$  is, the more the lattice is distorted. While at  $\Delta = 0.4$  and  $\Delta = 0.3$  open boxes still achieve long range order and high packing, at  $\Delta = 0.2$  the orientation defects destroy the lattice completely: no long range order is observed and a lower packing is reached.

We note that, the ot-tiling is reminiscent of the perfect kagome lattice observed in ref. 34, that is a hexagonal lattice with two kinds of pores: triangular and hexagonal ones. Both pore sizes are dependent on  $\Delta$ , with the pores becoming larger at more extreme ∆-values. We refer to such a tiling as open triangular-hexagonal tiling (oth-tiling) and we sketch it in Fig. 8c. To map the ot-tiling to the oth-tiling, the on-edge alignment of the unbonded edges - i.e., those edges that are not decorated with patches - must shift to an off-edge alignment. This shift would open up the hexagonal pores of the oth-tiling. Clearly, the ot-tiling prevails over the oth-tiling because hard

particles prefer arrangements that maximise edge-to-edge contacts. To stabilize a defect-free kagome lattice, neighboring boxes must be forced into an off-edge alignment. One strategy to reach this goal might involve the presence of patches in the outer perimeter of the open boxes in positions that stabilize the off-set edge contacts. The specific topology supporting the oth-tiling in ref. 34 can in fact be interpreted as a four patch extension of the ma-as topology that forms the open boxes.

Analogue to open boxes, we speculate that, if the star yield was high enough, an ot-tiling could be the assembly product of the 6-stars systems (see Fig. S5c in the ESI†). In this case, the distortion from such an hexagonal perfect lattice would be less extreme with respect to the open boxes as neighboring stars have no choice but to align parallel, thus leading to line defects (see Fig. S5d in the ESI†). Local hexagonal arrangements and line defects can be observed in simulations; however the hexagonal order does not prevail there as stars are not stable under compression. To enhance the stability of stars - and thus the quality of the second-stage assembly products - parallel bonds between the rhombi should be disfavored as it is this type of bonding that allows broken stars and disfavors chains (see Fig. S6a in the ESI†). A strategy to disfavor parallel bonds might consist in having different kinds of patches, where different types of patches attract each other (see Fig. S6b in the ESI†).

### Conflicts of interest

There are no conflicts to declare.

# Acknowledgements

CK and EB acknowledge support from the Austrian Science Fund (FWF) under Proj. No. Y-1163-N27. Computation time at the Vienna Scientific Cluster (VSC) is also gratefully acknowledged. VMD was used to generate the simulation snapshots. 44

#### Notes and references

- 1 W. Xu, Z. Li and Y. Yin, Small, 2018, 14, 1801083.
- 2 M. A. Boles, M. Engel and D. V. Talapin, Chem. Rev., 2016, **116**, 11220-11289.
- 3 R. Mérindol, E. Duguet and S. Ravaine, Chem. Asian J., 2019, 14, 3232-3239.
- 4 D. J. Kraft, R. Ni, F. Smallenburg, M. Hermes, K. Yoon, D. A. Weitz, A. van Blaaderen, J. Groenewold, M. Dijkstra and W. K. Kegel, Proc. Natl. Acad. Sci. U. S. A., 2012, 109, 10787-10792
- 5 S. Sacanna, D. J. Pine and G.-R. Yi, Soft Matter, 2013, 9,
- 6 B. Senyuk, Q. Liu, E. Bililign, P. D. Nystrom and I. I. Smalyukh, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2015, 91, 040501.
- 7 A. Azari, J. J. Crassous, A. M. Mihut, E. Bialik, P. Schurtenberger, J. Stenhammar and P. Linse, Langmuir, 2017, 33, 13834-13840.

Paper

8 Q. Chen, S. C. Bae and S. Granick, J. Am. Chem. Soc., 2012,

- 134, 11080-11083. 9 F. Lu, K. G. Yager, Y. Zhang, H. Xin and O. Gang, Nat.
- Commun., 2015, 6, 6912.
- 10 S.-M. Kang, C.-H. Choi, J. Kim, S.-J. Yeom, D. Lee, B. J. Park and C.-S. Lee, Soft Matter, 2016, 12, 5847-5853.
- 11 E. Bianchi, R. Blaak and C. N. Likos, Phys. Chem. Chem. Phys., 2011, 13, 6397.
- 12 A. Pawar and I. Kretzschmar, Macromol. Rapid Commun., 2010, 31, 150.
- 13 E. Bianchi, B. Capone, I. Coluzza, L. Rovigatti and P. D. J. van Oostrum, Phys. Chem. Chem. Phys., 2017, 19, 19847.
- 14 X. Ye, J. Chen, M. Engel, J. A. Millan, W. Li, L. Qi, G. Xing, J. E. Collins, C. R. Kagan, J. Li, S. C. Glotzer and C. B. Murray, Nat. Chem., 2013, 5, 466.
- 15 F. Ma, S. Wang, D. T. Wu and N. Wu, Proc. Natl. Acad. Sci. U. S. A., 2015, 112, 6307-6312.
- 16 K. Han, C. W. Shields IV, N. M. Diwakar, B. Bharti, G. P. Lòpez and O. D. Velev, Sci. Adv., 2017, 3, 1701108.
- 17 W. Li, D. Ruth, J. D. Gunton and J. M. Rickman, Chem. Phys., 2015, 142, 244705.
- 18 S. Whitelam, I. Tamblyn, J. P. Garrahan and P. H. Beton, Phys. Rev. Lett., 2015, 114, 115702.
- 19 M. Grünwald and P. L. Geissler, ACS Nano, 2014, 8, 5891-5897.
- 20 Q. Chen, S. C. Bae and S. Granick, J. Am. Chem. Soc., 2012, **134**, 11080-11083.
- 21 D. Morphew, J. Shaw, C. Avins and D. Chakrabarti, ACS Nano, 2018, 12, 2355-2364.
- 22 M. B. Zanjani, I. C. Jenkins, J. C. Crocker and T. Sinno, ACS Nano, 2016, 10, 11280-11289.
- 23 K. Aryana, J. B. Stahley, N. Parvez, K. Kim and M. B. Zanjani, Adv. Theory Simul., 2019, 2, 1800198.
- 24 M. N. O'Brien, M. R. Jones, B. Lee and C. A. Mirkin, Nat. Mater., 2015, 14, 833-839.
- 25 M. N. O'Brien, M. Girard, H.-X. Lin, J. A. Millan, B. L. Monica Olvera de la Cruz and C. A. Mirkin, Proc. Natl. Acad. Sci. U. S. A., 2016, 113, 10485-10490.

- 26 W. Liu, J. Halverson, Y. Tian, A. V. Tkachenko and O. Gang, Nat. Chem., 2016, 8, 867-873.
- 27 W. Liu, M. Tagawa, H. L. Xin, T. Wang, H. Emamy, H. Li, K. G. Yager, F. W. Starr, A. V. Tkachenko and O. Gang, Science, 2016, 351, 582-586.
- 28 S. Ithurria, M. D. Tessier, B. Mahler, R. P. S. M. Lobo, B. Dubertret and A. L. Efros, Nat. Mater., 2011, 10, 936-941.
- 29 A. C. Stelson, W. A. Britton and C. M. L. Watson, J. Appl. Phys., 2017, 121, 023101.
- 30 P. W. K. Rothemund, Nature, 2006, 440, 297-302.
- 31 G. Tikhomirov, P. Petersen and L. Qian, J. Am. Chem. Soc., 2018, 140, 17361-17364.
- 32 X. He, Y. He, M.-S. Hsiao, R. L. Harniman, S. Pearce, M. A. Winnik and O. Manners, J. Am. Chem. Soc., 2017, 139, 9221-9228.
- 33 G. Chen, G. Zhang, B. Jin, M. Luo, Y. Luo, S. Aya and X. Li, I. Am. Chem. Soc., 2019, 141, 15498-15503.
- 34 C. Karner, C. Dellago and E. Bianchi, Nano Lett., 2019, 19, 7806-7815.
- 35 N. Pakalidou, J. Mu, A. J. Masters and C. Avendaño, Mol. Syst. Des. Eng., 2020, 5, 376-384.
- 36 Ł. Baran and W. Rżysko, Mol. Syst. Des. Eng., 2020, DOI: 10.1039/C9ME00122K.
- 37 E. G. Golshtein and N. Tretyakov, Modified Lagrangians and monotone maps in optimization, Wiley, 1996.
- 38 M. C. L. Gottschalk and C. Manocha, Comput. Graph., 1996, 30, 171-180.
- 39 S. Whitelam, I. Tamblyn, P. H. Beton and J. P. Garrahan, Phys. Rev. Lett., 2012, 108, 035702.
- 40 S. Whitelam and P. Geissler, J. Chem. Phys., 2007, 127, 154101.
- 41 S. Whitelam, Mol. Simul., 2011, 37, 606-612.
- 42 C. Karner, C. Dellago and E. Bianchi, J. Phys.: Condens. Matter, 2020, 32, 204001.
- 43 E. P. Bernard and W. Krauth, Phys. Rev. Lett., 2011, 107, 155704.
- 44 W. Humphrey, A. Dalke and K. Schulten, J. Mol. Graphics, 1996, 14, 33-38.