Self-assembly of colloidal superballs under spherical confinement of a drying droplet

Citation for published version (APA):

Document license:
CC BY-NC-ND

DOI:
10.1016/j.jciso.2021.100037

Document status and date:
Published: 01/04/2022

Document Version:
Publisher’s PDF, also known as Version of Record (includes final page, issue and volume numbers)

Please check the document version of this publication:

• A submitted manuscript is the version of the article upon submission and before peer-review. There can be important differences between the submitted version and the official published version of record. People interested in the research are advised to contact the author for the final version of the publication, or visit the DOI to the publisher’s website.
• The final author version and the galley proof are versions of the publication after peer review.
• The final published version features the final layout of the paper including the volume, issue and page numbers.

Link to publication

General rights
Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.
• Users may download and print one copy of any publication from the public portal for the purpose of private study or research.
• You may not further distribute the material or use it for any profit-making activity or commercial gain
• You may freely distribute the URL identifying the publication in the public portal.

If the publication is distributed under the terms of Article 25fa of the Dutch Copyright Act, indicated by the “Taverne” license above, please follow below link for the End User Agreement:
www.tue.nl/taverne

Take down policy
If you believe that this document breaches copyright please contact us at:
openaccess@tue.nl
providing details and we will investigate your claim.

Download date: 12. Jun. 2022
Self-assembly of colloidal superballs under spherical confinement of a drying droplet

Sarah Schycka,², Janne-Mieke Meijerb,², Lucia Baldaufc,¹, Peter Schallc, Andrei V. Petukhovd,e, Laura Rossia,*

a Delft University of Technology, Department of Chemical Engineering, 2629 HZ, Delft, the Netherlands
b Eindhoven University of Technology, Department of Applied Physics and Institute for Complex Molecular Systems, Eindhoven, the Netherlands
c University of Amsterdam, Institute of Physics, Science Park 904, Amsterdam, the Netherlands
d Utrecht University, Debye Institute for Nanomaterials Science, Utrecht, the Netherlands
e Eindhoven University of Technology, Department of Chemical Engineering and Chemistry, Eindhoven, the Netherlands

A R T I C L E   I N F O

Keywords:
Small angle X-ray scattering
Colloidal superballs
Core-shell particles
Spherical confinement

A B S T R A C T

Understanding the relationship between colloidal building block shape and self-assembled material structure is important for the development of novel materials by self-assembly. In this regard, colloidal superballs are unique building blocks because their shape can smoothly transition between spherical and cubic. Assembly of colloidal superballs under spherical confinement results in macroscopic clusters with ordered internal structure. By utilizing Small Angle X-Ray Scattering (SAXS), we probe the internal structure of colloidal superball dispersion droplets during confinement. We observe and identify four distinct drying regimes that arise during compression via evaporating droplets, and we track the development of the assembled macrostructure. As the superballs assemble, we found that they arrange into the predicted paracrystalline, rhombohedral C1-lattice that varies by the constituent superballs’ shape. This provides insights in the behavior between confinement and particle shape that can be applied in the development of new functional materials.

The self-assembly of colloidal particles is an ideal mechanism for structuring matter at the nano- and microscale, and can produce materials of interest for different applications, such as photonics, optoelectronics, catalysis or energy storage [1,2]. To direct the self-assembly process towards the desired structures, different methods have been developed and explored [3]. One unique way to tune the spontaneously formed structures is to assemble the colloidal particles (of sizes from 2 nm to 1 μm) under spherical confinement induced by evaporating micro-emulsion droplets or larger, millimeter-sized dispersion droplets [4–7]. Colloidal spheres typically crystallize into face centered cubic (FCC) crystal phases. When this type of crystallization occurs under spherical confinement, unique optical properties can arise, i.e., structural colors, in the final clusters. In these cases, cluster diameters can range from <50 μm [8] to the millimeter-scale [9,10]. Furthermore, control over the number of particles inside a dispersion droplet has been shown to produce novel structures, such as icosahedral clusters [11,12] and magic number clusters [13]. Depending on the initial colloidal volume percent and the drying substrate properties, dispersion droplets drying on superhydrophobic surfaces can form various cluster shapes, e.g., doughnuts or flattened discs [10,14] and spheres [4,9].

Besides finely tuning the assembly process, improvements in the synthesis of anisotropic colloidal particles have opened the path for more structurally varied assemblies [15]. Consequently, there is a great diversity within the colloidal assemblies of different anisotropic and polyhedral shapes [16–18]. Combining spherical confinement with anisotropic shapes can lead to even more complex micro structures. For instance, simulations of different polyhedral shapes showed that spherical confinement leads to unique clusters distinctly different from the packings of spheres [19]. For rod-like colloids, it was found that their disordered so-called supraparticles possess unique scattering enhancing properties in contrast to their spherical counterparts [20].

Among available anisotropic particles, the superball shape, which encompasses the transition from a sphere to a cube via a rounded cube,
has become of interest due to its unique shape-dependent phase behaviour [21–23] and experimental availability [24,25,26]. For hard superballs, the densest packing is a rhombohedral-like crystal structure that, with increasing asphericity of the superballs, continuously evolves from an FCC lattice (spheres) to a simple cubic (SC) lattice (perfect cubes) [21, 22,24,25,27]. Before assembling into their densest packings from a liquid phase, superballs can form a plastic crystal (or rotator) phase which has translational order and rotational mobility [23,27]. Experimentally, the phase diagram of micron-sized superballs has been studied via crystallization under gravitational settling in a capillary [27] and depletion-assisted assembly [26,28]. For nano-sized superballs, the dense packings have also been investigated [24,25]. These studies also confirmed the formation of dense rhombohedral-like structures of which the exact packing is dependent on the asphericity of the superball shape.

In addition, the self-assembly of micron-sized superballs into monolayers driven by solvent evaporation processes on flat substrates was investigated [39,40]. These studies showed that solvent flow and immersion capillary forces, which can reach 10^4 kT for micron-sized colloids, will lead to the formation of similar dense rhombohedral-like crystal structures but also revealed that the out-of-equilibrium process induces the formation of lower density crystal lattices with hexagonal-like structures as well as disordered packings.

So far, only a few studies have investigated the effect of spherical confinement on the assembly of superball (or cubic) particles and the relationship between particle asphericity and the resulting densely packed structure symmetries. For superball nanoparticles with increasing asphericity, the confined structure changes from icosahedral clusters with strong local orientation to simple-cubic structures [31]. Cubic nanoparticles, with very sharp edges, assemble at the surface and in bulk of the droplet and form densely packed clusters with random orientations throughout the confined structure [32]. Moreover, it has been shown that when nanocubes form dense supraparticles, the finite corner roundness and the surface tension of the confining droplet determine the superstructure morphology [33]. To understand the final self-assembled supraparticles’ structure formed by superballs, it is crucial to understand the dynamics of the particles during droplet evaporation. For this, small angle x-ray scattering (SAXS) is a very powerful tool that allows visualization of the assembly process of different nanoparticles during spherical confinement in detail [32,34–36] and has been used to resolve micron-sized superball particle assembly [27].

In this work, we use in-situ synchrotron small angle X-ray scattering with microradian resolution (μrad-SAXS) to reveal the crystallization process of micron-sized silica superball shells under spherical confinement through droplet drying and elucidate the structural transitions during assembly. We are able to follow the droplet evaporation process in detail and identify different stages during the process. As the solvent evaporates, the particle concentration increases. After this stage, dewetting of the superball shells occurs before full solvent evaporation. We find that crystalline structures start to form as particle concentration increases, and that the structures remain throughout the later stages. In addition, we find that as we increase the asphericity of the superballs, the assembled supraparticles’ structures are polycrystalline assemblies of their predicted rhombohedral-like lattices.

1. Results

To study the self-assembly process of superball particles, we synthesized four samples of hollow silica superballs. We note that superballs are a family of shapes defined by \( \frac{x^2}{a^2} + \frac{y^2}{a^2} + \frac{z^2}{a^2} \geq 1 \) where \( x, y \) and \( z \) are the unit axis pointing toward the superball faces, \( a \) is \( \frac{1}{3} \) the distance between two opposite faces, and \( m \) is the shape parameter [21,37]. To create superballs with a range of \( m \) values, we coat hematite superball seeds in silica where increasing the silica shell thickness, \( t \), decreases the resulting \( m \) values [28]. The hematite seeds are etched away through the porous silica superball shell (Fig. 1A). For example, \( m = 2.6 \) particles have shells of \( t = 120 \) nm and particle sizes of \( 2 \cdot a = 1240 \) nm with a size dispersity of 4%. A table of all sample parameters is provided in Supplementary Table S1. For particles with \( m = 2 \) (spheres), we grew two sets of silica particles without a template which resulted in solid silica spheres with radii of \( r = 206 \) nm and \( r = 233 \) nm [38].

We fabricate assemblies of colloidal superballs by drying 1.5 μL dispersion droplets of 15 vol% superballs in MiliQ water on the surface of a superhydrophobic substrate. We add small pinning positions to the surface of the substrate with a needle tip to prevent droplet rolling. These types of dispersion droplets dry via diffusion of the water vapor into the air [14].

To investigate how the structure assembles under spherical confinement, we performed in-situ SAXS measurements on a series of drying dispersion droplets where each contained different \( m \) valued superballs, as schematically shown in Fig. 1B. To control the evaporation rate throughout experiments, droplets were dried at ~64% relative humidity in a custom-made chamber (Supplementary Fig. S1). As water evaporates, the droplet contracts from the initial diameter of 1.5–2 mm to ~1 mm (Supplementary Fig. S2) and eventually yields a closed-packed assembly which is apparent from the rise in iridescent colors resulting from the Bragg scattering of light as seen in Fig. 1C [6,9]. The synchrotron X-ray beam, of width 500 μm, passed through approximately the center of each droplet, and 2D images were recorded every ~4 seconds by the high-resolution 2D X-ray detector for 30–35 min as the droplet dried.

![Fig. 1.](image-url) (A) Transmission electron microscopy images of a solid silica sphere and hollow silica superballs with different shape parameters, \( m \), shell thicknesses, \( t \), and radii, \( a \). Scale bars are 200 nm. (B) Schematic representation of the in-situ SAXS setup. A dispersion droplet on a superhydrophobic surface is placed inside a humidity chamber. Scattered x-rays are collected on a 2D detector with a wedge-shaped beam-stop. Schematic is not to scale. (C) A time sequence of 42 images of a solid silica sphere and hollow silica superballs with different size parameters. (D) A table of all sample parameters is provided in Supplementary Table S1. For particles with \( m = 2.6 \) (spheres), we grew two sets of silica particles without a template which resulted in solid silica spheres with radii of \( r = 206 \) nm and \( r = 233 \) nm [38]. (E) A schematic representation of the different observed stages of drying: Dilute (non-interacting), structure formation, dewetting, and the final dried assembly.
From the 2D images, the 1D radially averaged scattering profiles were extracted. A typical set of scattering profiles collected in time during the droplet drying process for superballs, with $m = 2.6$, is displayed in Fig. 1D. Time resolved scattering profiles for all investigated droplets are included in Supplementary Figs. S3 and S4.

Generally, the scattering intensity, $I(q)$, from a collection of discrete particles is dependent on the form factor, $P(q)$, and the structure factor, $S(q)$. $P(q)$ arises from the particles' shape and size, while $S(q)$ arises from the particles' organization. The total scattering intensity, $I(q)$, for superball particles can be approximated by the $I(q)$ for monodisperse spherical particles [27,39]. In this approximation, $I(q) \propto \phi_{part} \cdot P(q) \cdot S(q)$ where $\phi_{part}$ is the particle volume fraction [32]. In addition, the overall droplet drying process can be monitored via the normalized scattering power, $Q^*$, which is defined by $Q^* = Q/Q_{\text{max}} \propto \phi_0(1 - \phi_{SB})$ where $Q = \int_0^\infty q^2 \cdot I(q) dq$ is the Porod invariant, $Q_{\text{max}}$ is the maximum $Q$ value, $\phi_0$ is the volume fraction of superballs, and $\phi_{SB}$ is the scattering length density (SLD) which arises from the difference in electron density between the particle and surrounding medium [34].

Throughout the droplet drying process, we observe four distinct regimes in the $I(q)$ patterns as shown schematically in Fig. 1E. Initially, the superball particles in the droplet are in a dilute, non-interacting, fluid state where we can assume that $S(q) \approx 1$ and that $P(q)$ dominates the scattering profile. We note that we assume $S(q) \approx 1$ in the observed experimental $q$ region due to the lack of additional long range effects from the superballs [40] because we consider the superballs to be hard particles. We check this assumption via the Percus-Yevick equation [41].

Fig. 1D. Time resolved scattering profiles for droplets with $m = 2.6$. Dashed lines show the peak position ratios over time for the superballs. $\phi_0 = 15$ vol % (Supplementary Fig. S5). During the first drying period, the evaporation and confinement do not influence the system, which is evident from the limited changes in the patterns and the little variation in $Q^*$. As water evaporates from the droplet, the confining volume decreases, and the concentration of particles increases. As a result, inter-particle interactions arise, leading to $S(q) \neq 1$, and $S(q)$ peaks start to appear in the $I(q)$ profiles indicating structure formation. Near the final stage of drying, dewetting occurs where the remaining water inside and outside the particles evaporates, and the surrounding medium of the particles becomes air. This change of medium from water to air dramatically increases the difference between SLDs of the colloids and the surrounding medium, $\Delta \rho$. This step in the drying process leads to a change in the overall intensity of the 1D profiles and appears as a rapid increase in $Q^*$. Therefore, we utilize the inflection point of $Q^*$ to determine the beginning of dewetting (see inset in Fig. 1D). For all investigated superball droplets and their droplets, dewetting began at similar times (at $t = 24.1 \text{ min}$ for $m = 2.6$, at $21.7 \text{ min}$ for $m = 2.8$, $3.1$, and $3.3$, and at $22.9 \text{ min}$ for both $m = 2$ containing samples). Discrepancies between dewetting times are likely due to the 2-5 min time variation in placing the droplet and beginning the SAXS measurements. Eventually, all of the remaining water in the system evaporates and only the dried assembly of particles remains. When the dried assembly is formed, the 1D profiles and $Q^*$ plateau. During the SAXS measurements, not all investigated droplets formed a dried assembly (Supplementary Figs. S3 and S4).

Before characterizing the full drying process, we first characterize the particles in the initial, dilute state where $P(q)$ dominates the scattering profile. $P(q)$ was determined to be the initial free particle scattering profiles when $t = 0$, $I_{\text{free}}(q)$ and is shown for all investigated particle shapes with $2 \leq m \leq 3.3$ in Fig. 2. As is well known for hollow spherical shells, the $P(q)$ of a hollow object will be dependent on the details of the shell thickness and the contrast between the solvent, shell, and core [41]. In the case of hollow superballs, the $P(q)$ is determined by the superball shell thicknesses ($t$), inner shape ($t_{\text{inner}}$), and outer shape ($t_{\text{outer}}$) parameters [27]. The slight variations visible between the $P(q)$ of the different hollow superballs seen in Fig. 2 can be attributed due to these differing parameters. In addition, it is clear that the $P(q)$ minima are not very distinct, which we attributed to a small size dispersity ($3 - 5\%$) of the particles [42]. It is important to note, for drying hollow superballs, using $I_{\text{free}}(q)$ as $P(q)$ is only valid when all the particles' cores match the surrounding medium’s electron density. When dewetting occurs, particle cores do not empty at the same time and $P(q) \neq I_{\text{free}}(q)$ as shown schematically in Fig. 1E. Additionally, $S(q)$ changes as neighboring particles no longer possess the same SLD (for more details see Supplementary Fig. S6).

We can visualize the drying process by examining the 2D scattering images as done in Fig. 3A for a drying droplet with $m = 2.6$ superballs. Here, we observe isotropic scattering profiles from the fluid until Bragg peaks appear in the first ring after 10 min, indicating the formation of crystalline structures. These $S(q)$ peaks are present through the dewetting point $(t = 24.1 \text{ min})$. Compared to distinct single crystalline 2D SAXS patterns observed for 3D superball sediments [27,43], the radial width of these peaks is larger and the fact that higher order peaks are absent indicate that the resultant structure is polycrystalline where the domains have only weak orientational correlations with each other. For all investigated superball dispersion droplets, we find similar scattering patterns. Although as $m$ increases, the peaks broaden, and for $m = 3.3$ only a ring can be seen (see Supplementary Fig. S8).

To further understand the structure formation, we examine the 1D $S(q)$ curves. We can extract $S(q)$ by dividing $I(q)$ with the $P(q) = I_{\text{free}}(q)$ up to dewetting phase where the patterns start to change dramatically (as discussed above). Fig. 3B shows selected $S(q)$ curves for superballs with $m = 2.6$ up to $t = 24.1 \text{ min}$. It can be seen that three consistent, albeit broad, peaks develop in the curves around $t = 10 \text{ min}$ indicating that structure formation has begun. During drying, all peak positions continuously shift towards higher $q$ values demonstrating decreasing inter-particle distances between the superball particles (Supplementary Fig. S7). After $t = 20 \text{ min}$, a shoulder emerges from the $q_0$ peak.

To reveal how the structure evolves, we follow the ratio between the peak positions and the first peak, $q_0/q_0$ [24,27]. Peak positions and widths were determined by fitting the peaks with a Voigt function [44]. Fig. 3C shows the peak position ratios over time for the $m = 2.6$ superball droplet. The ratio remains relatively constant throughout droplet drying and suggests that no significant phase transitions occur. From the $q_0$ peak width, the average crystallite size can be calculated using the Scherrer equation [45]. To easily compare the result between different cubes, the average crystallite size, $D$, is normalized by dividing the superball side length, $(2 - a)$, to find $D' = D/(2 - a)$. Fig. 3D shows that $D'$ begins to increase.
rapidly after 16 min as the droplet dries. This indicates that the average crystalline domain grows.

To understand the influence of particle shape during assembly, we explore how $S(q)$ evolves for all dispersion droplets of superballs with $m \leq 3.3$. Fig. 4A–C shows the temporal $S(q)$ curves for $m = 2.8$, $m = 3.1$, and $m = 3.3$. In all droplets, we observe a trend similar to $m = 2.6$ where $S(q) > 1$ at $t \approx 7$ min. The peak positions shift to higher $q$ values, and a shoulder emerges from the $q_0$ peak. To confirm that the structure remains after dewetting occurs, we can extract $S(q)$ of a dried assembly and compare to the structure before dewetting by rescaling $P(q)$ [41]. Only particles with $m = 3.3$ formed a completely dried assembly which was determined by a plateau at the end of $Q^*$ (Supplementary Figs. S3–S4). The solid black curve in Fig. 4D depicts the structure of the dried assembly and is compared to $S(q)$ before dewetting (dashed black curve). The increase in intensity is due to the change in surrounding medium [41]. Otherwise, there are no significant changes to $S(q)$ after dewetting.

Furthermore, we compare the average crystallite size, $D^*$, over time for all investigated droplets in Fig. 4E. As the droplets dry, we observe that the average crystallite size increases for all investigated droplets. Indeed, the average crystallite size at the onset of structure formation ($\approx 10$ minutes) spans approximately 5 particle diameters for all droplets. At 22 min before dewetting occurs, $D^* \approx 5$ particles per crystallite for $m = 2.6$ and $m = 2.8$. Then, $D^*$ increases with the $m$ value where the most cubic-like particle assembly ($m = 3.3$) has $D^* \approx 9.8$ particles per crystallite.

To determine the crystal structures that have formed in the droplets, we first extract $S(q)$ for the spherical particle dispersion droplets just before dewetting (Fig. 5A). To compare different dispersions, we re-scaled $q$ of all the $S(q)$ curves by the corresponding $q_0$ position of the first $S(q)$ peak. In general, spherical colloids do not assemble into pure FCC lattices, but they form a random hexagonal close packed (RCHP) structure that is a mixture of both the FCC and hexagonal close packed (HCP) lattices [46]. Therefore, we compare the $S(q)$ curves from two separate droplets with different solid silica sphere particle radii ($R = 206, 233$ nm) to the expected $S(q)$ peaks for FCC (black model lines) and HCP (green lines) in Fig. 5A. We find that the spherical assemblies produced similar broad peaks near expected FCC peak positions for the $hkl$ planes $111, 311, 331, \text{and } 440$.

On the other hand, as the $m$ value increases, we expect that perfect cubic particles pack into simple cubic (SC) lattices [32]. Therefore, in Fig. 5B, we compare the curves to the expected $S(q)$ peaks for FCC (black model lines) and SC lattices (gray model lines). For all $m$, it is clear that the peaks do not perfectly match either the FCC peaks or the SC peaks. Instead as we increase $m$, we observe a slight shift of peak positions away from $q_0$.

Fig. 3. Structural evolution for a drying droplet containing superballs with $m = 2.6$. (A) Selected 2D SAXS patterns showing the rise of Bragg peaks during drying. (B) Selected $S(q)$ curves extracted until the dewetting phase. (C) The ratio of peak positions to the first peak location ($q/q_0$) at selected times. (D) The average crystallite size normalized to the particle size ($2 \cdot a$) over droplet drying.

Fig. 4. Structural evolution of superball dispersion droplets during drying. Selected $S(q)$ curves extracted until the dewetting phase for (A) $m = 2.8$, (B) $m = 3.1$, and (C) $m = 3.3$ with a time-step of 1.2 min between curves. (D) Comparison of $S(q)$ curves from before dewetting (black, dashed) to the final dried assembly (black, solid) for $m = 3.3$. (E) The normalized average crystallite size for all superball systems at 10 min (blue bars) and 22 min (orange bars). (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)
Indeed, simulations and experiments have shown that the equilibrium structure formed by superballs, with $m \geq 2.30$, is a densely packed rhombohedral crystal lattice, described by the C1-lattice [21–23,27]. The C1-lattice is dependent on the superball $m$ parameter and describes a continuous transition from an FCC structure for $m = 2$ to an SC structure for $m \to \infty$ as shown in Fig. 5C.

To compare our assembled structures to the C1-lattice, we calculate how the $S(q)$ peaks evolve as the $m$ values increases. The lattice vectors for the C1-lattice are defined by Jiao et al. [21] as $e_1 = -2(s + 2^{-1/m})i + 2j + 2k$, $e_2 = -2s + 2j + 2s + 2^{-1/m}j$, and $e_3 = -2i + 2s + 2^{-1/m}j + 2sk$ where $s$ is the smallest positive root of the equation $(s + 2^{-1/m})^m + 2^{-m} = 1$. The structure varies as $m$ increases by changing the angle, $α$, between lattice vectors: $α = \arccos \frac{\sum_{i,j} e_i e_j}{m}$, where $i,j = 1, 2, 3 (i \neq j)$. It has been shown that when superballs assemble into their densest packing, we can directly map their expected angle, $α$, from the experimental $m$ value [27]. We note that as $m$ increases from 2 to 3.3, the angle, $α$, increases from 60° to 69° (Supplementary Fig. S11). Achieving higher angles ($α > 80^\circ$) requires particles with high $m$ values ($m > 10$).

Then, we can follow the transition between FCC and SC lattices of the C1 lattice via the ratio of the diffraction peak position for 60° < $α$ < 90°. This ratio is defined as

$$\frac{q_{\text{SC}}}{q_{\text{FCC}}} = \frac{(h^2 + k^2 + l^2)\sin \alpha^2}{\sin \alpha^2} + \frac{2(hk + kl + lh)(\cos \alpha^2 - \cos \alpha)}{\sin \alpha^2}$$

where $q_{\text{SC}}$, $q_{\text{FCC}}$ are the peak positions of allowed (hkl) reflections. Fig. 5D shows how the ratios of several hkl peak positions are expected evolve according to eq. (1) for angles from 60° to $α \leq 90$ (lines). The respective FCC and SC hkl values related to the calculated rhombohedral planes are included. Additionally, we overlay the experimental data points (symbols) on top of the lines for comparison. As $α$ increases, the diffraction ratio generally grows for most hkl planes. Higher $q$ value peaks tend to increase more rapidly compared to lower $q$ value peaks. However, that is not the case for all hkl planes such as the horizontal lines associated to the transitions into the SC h00 planes. While the experimental values are only available up to $m = 3.3$, corresponding to $α \approx 70^\circ$, the peak ratio appears to generally follow the predicted trends. We observe that the emerging peak off of $q_0$ closely follows the FCC 200 → 110 SC plane transition. Similarly, the higher-order peaks also increase with increasing $m$.

2. Discussion

Our μrad-SAXS results show that the spherical confinement, due to drying dispersion droplets of different superball colloids, leads to the formation of polycrystalline assemblies with their expected rhombohedral C1-lattice packing [21]. We further find that the average crystalline domain increases with the particles’ $m$ value. Our findings are in part in agreement with the results of Wang et al. [31] who showed that for small droplets, with $\approx 2000$ cubic nanoparticles, there are short range correlations between the particle positions of dried superball assemblies with low $m$ values, and long-range correlation increases with increasing $m$. However, in contrast to their study, we do not find the formation of icosahedral clusters for low $m$ as the droplets investigated here are much larger and contain $\approx 2 \times 10^5$ superball particles. In this respect, our results are more in line with studies focusing on droplets with a larger number of particles. For nanocubes with $m > 5$, Aghe et al. [32] showed that spherical confinement in a levitating drop leads to polycrystalline domains where domains at the surface were aligned to the droplet interface. For micron-sized spheres, Marin et al. [4] showed that droplets...
on superhydrophobic surfaces, at high enough particle concentration, will form densely packed structures with many different crystalline domains also aligned to the droplet surface. The reason for this polycrystalline structure is that densification of a droplet leads to multiple nucleation sites when the droplet contains a large number of particles. Interestingly, we did observe the appearance of distinct six-fold Bragg peaks on top of the $S(q)$ ring in the 2$d$ rad-SAXS patterns of dispersion droplets with $m < 3.3$. This could indicate that while a large part of the surface crystallites are randomly oriented, the center part of the droplet may contain a structure with long-range order, but more investigations are needed to confirm this.

By following the full drying process, we showed that the system densifies and the crystalline structure starts to form as the droplet evaporates. Interestingly, the forming crystal structures did not undergo any phase transitions, which might be expected based on a previous study by Meijer et al. [27] who showed that superball colloids, upon slow sedimentation, will first form a rotator FCC crystal before transforming into their densest rhombohedral packing. Clearly, the current assembly process during spherical confinement occurs via different pathways. We further revealed that a dewetting stage occurs, during which the water evaporates from between and inside the hollow colloids, leading to the dramatic change in the particle $P(q)$. Since the water does not evaporate from all of the porous hollow colloids at one instant, we do not observe such a significant change for the solid spheres, and this has not been observed for other droplet drying studies of solid particles as well. Thus these hollow particles allowed us to obtain more insight into the local evaporation process as we tracked the changing intensity in $P(q)$ and the corresponding $I(q)$. In addition, strong capillary forces acting on the colloidal particles during dewetting of the droplet, which can reach $10^6$ kT [47], do not lead to rearrangement of the crystalline structures that have formed, as the $S(q)$ before and after dewetting are the same.

In conclusion, we investigated the structural evolution of colloidal superball assemblies during spherical confinement induced by a drying dispersion droplet. The presented results show that the structure of the assembled superballs nucleate into the predicted C1-lattice. As the droplet evaporates, we found that the structure does not undergo any additional phase transitions throughout drying. The resulting assembly contains several small crystallites that increase in size as the superballs become more cubic-like indicating a polycrystalline structure, and the assembly exhibits optical reflections on its surface.

As our understanding of superball assemblies continues to advance, we anticipate that insights into the relationships between the constituent anisotropic particles and resulting assembly’s structural color and mechanical properties will arise. It has been shown that by switching from spherical to rod-like particles, dramatic changes in the assembly’s structural color can be obtained after spherical confinement [20]. Similarly, particle size and interparticle bonds influence the mechanical properties of the assembly [48]. Looking forward, we anticipate that by retaining the magnetic hematite seed inside the silica superball [43], we can explore how the additional magnetic dipole interactions influence the assembly under spherical confinement.

3. Methods

3.1. Particle preparation

Hematite cube template particles were prepared following a sol-gel procedure [49]. Silica shells were grown following Ref. [50]. The hematite templates were dissolved through the addition of hydrochloric acid to produce hollow shells. The complete particle synthesis procedure to obtain different m values can be found in Ref. [38]. Silica spheres were prepared using a variation on the Stober method [51] from Ref. [38] to create spheres with diameter of $D \approx 500$ nm. The obtained particles were characterized by transmission electron microscopy (TEM - Philips TECNAI12/20). Details on the properties of all prepared particles are presented in Supplementary Table S1.

3.2. Assembly process

3.2.1. Superhydrophobic substrate

Superhydrophobic substrates with contact angles above $160^\circ$ were prepared following a modified procedure from Deng et al. [52]. Briefly, a thin layer of soot from the flame of a paraffin candle is deposited on a cleaned glass substrate. Then, the substrate is coated with a thin silica shell by enclosing it in a desiccator with two open containers of 2 mL tetraethyl orthosilicate and 2 mL NH$_3$ (29 wt%) for 24 h. The substrate is then calcinated in an Even heat Rampmaster RM2 furnace by heating at $600 \, ^\circ\text{C}$ for 2 h. After, $350 \, \mu\text{L}$ of 1% trichloroethylene in chloroform is dropcasted on the substrate and allowed to evaporate. Droplet pinning positions are created on the substrate via notches on the surface.

3.2.2. Spherical confinement of colloids

Self assembled macrostructures of colloidal spheres and superballs were formed by depositing and drying 2–5 μL droplets of colloidal dispersions from 5 to 30 vol% on a superhydrophobic substrate at the pinning positions. These are necessary to prevent the droplets from moving and escaping the superhydrophobic surface. Evaporation rates can be controlled via drying inside a humidity chamber (See Supplementary Fig. S1).

3.2.3. Small angle X-ray scattering (SAXS) measurements

rad-SAXS measurements were conducted at the beamline BM26B DUBBLE [53] at the European Synchrotron Radiation Facility (ESRF) in Grenoble, France using a microradian setup [54] employing compound refractive lenses [55]. The incident X-ray beam with a photon energy of 13 keV ($\gamma = 1.033\,2\text{A}$) was focused on a Photonic Science CCD detector with 4,008 $\times$ 2,671 pixels and a pixel size of $22 \times 22\, \mu\text{m}$ at a distance of 7.11 m. The detector was protected from the direct beam by a wedge-shaped beam-stop. Substrates with the drying droplets were placed horizontally in the X-ray beam. Measurements were performed during the full evaporation process at the position of the initial center of the drying droplet.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgements

We thank the Netherlands Organization for Scientific Research (NWO) for the provided beam-time. NWO is also acknowledged for financial support of L.R. through a VENI grant(680-47-446) and J.M.M. (016.Veni.192.119). L.B. acknowledges the support by a Studienstiftung des Deutschen Volkes research grant. We are also grateful to the DUBBLE personnel of the European Synchrotron Radiation Facility in Grenoble for assistance with the small angle X-ray scattering experiments.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jcis.2021.100037.

References


[27] Stoeb Werner, Arthur Fink, Ernst Bohn, Controlled growth of monodisperse silica spheres in the micron size range, J. Colloid Interface Sci. 21 (1) (1968) 62–69.

