

Modularly Constructed Polyhedral Oligomeric Silsesquioxane-Based Giant Molecules for Unconventional Nanostructure Fabrication

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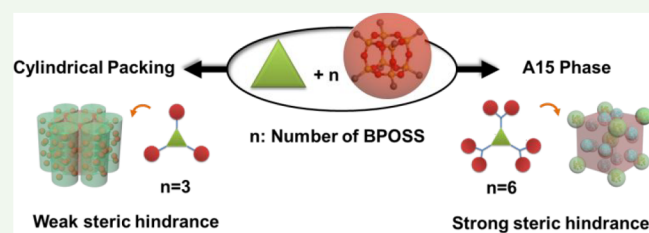
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ABSTRACT: Controlled assembly of nanoscale building blocks is a promising approach to obtain functional materials with unique properties. Here, we report a way to manipulate the supramolecular structures of giant molecules based on discotic triangle cores and isobutyl polyhedral oligomeric silsesquioxanes (BPOSS) nanoparticles (NPs). It is found that depending upon the number of BPOSS at the periphery of the discotic cores, the packing of these nanoscale components (discotic core and POSS) could be manipulated into either cylindrical or Frank–Kasper (F–K) A15 ($Pm\bar{3}n$) phases. The formation of these supramolecular nanostructures is mandated by the balance between the stacking of the discotic cores and the steric hindrance effect of the BPOSS NPs. This strategy to manipulate the packing of nanoscale building blocks for different supramolecular nanostructures including the fabrication of cylindrical structures and A15 ($Pm\bar{3}n$) phases may be extended to other nanoscale building blocks for future development of materials with complex structures as well as tailored functionalities and properties.

KEYWORDS: nanoparticles, Frank–Kasper phase, self-assembly, supramolecular chemistry, POSS



1. INTRODUCTION

Achieving materials with desired properties and functionalities via reverse thinking and designing has been demonstrated as one of the most promising ways in self-assembly (“bottom-up”) approaches. In the past two decades, different functionalized nanoscale building blocks have been used to achieve various interesting nanostructures and properties in soft matters.^{1–5} Specifically, Frank–Kasper (F–K) phases, which originally appeared in metal-alloys with specifically required spherical motifs,^{6,7} are receiving researchers’ attention since they have been observed in soft materials, such as supramolecular dendrimers,^{8–12} self-organizable dendronized polymers,^{13–16} block copolymers,^{17–19} surfactants,^{20–23} and giant molecules.^{24–26} For example, in 1997, the first thermotropic A15 ($Pm\bar{3}n$) phase in soft matter was discovered in dendrimers by Percec et al.²⁷ Since then, a number of dendrimers were found to assemble into the A15 ($Pm\bar{3}n$) phase,^{27–32} σ ($P4_2/mnm$) phase,³³ and quasicrystal phase^{34,35} together with the traditional phases, providing a “nanoperiodic table” of supramolecular structures.^{10,12,36} Bates et al. discovered the F–K σ ($P4_2/mnm$) phase, C14 ($P6_3/mmc$) phase, and C15 ($Fd\bar{3}$) phase in sphere-forming block copolymer melts.^{17–19} Mahanthappa et al. found F–K phases formation in surfactant micelles.^{20–23} In the meantime, some simulation and theoretical works about F–K phases have been carried out.^{37–46} For example, Kamien et al. investigated the theory of F–K phase formation using a packing model of a hard core

and soft corona system.^{41,42} Goddard et al. conducted molecular dynamic simulation of supramolecular dendrimer balls on forming A15 ($Pm\bar{3}n$) structures.³⁷ Glotzer et al. studied the simulation on self-assembly of soft matter, including conventional structures as well as quasicrystals and their approximants.^{38–40}

According to those simulations, the building block units must possess specific topological restrictions and secondary interactions. The building blocks in the previously reported examples are mostly soft. Recently, the F–K phases have also been discovered in several categories of giant molecules.^{24–26} Giant molecules are a kind of nanoscale macromolecules constructed by the precise molecular nanoparticles (MNPs) as composition units, such as POSS, fullerenes, polyoxometalates, etc.^{47–49} The term “giant” is referring to comparison with their small molecular counterparts as the MNP units resemble atoms but are thousands of times larger. Similar concept, such as the nanoelements, have also been used.^{12,36} Due to the shape and volume-persistence of the MNPs, the principles of the self-assembly of giant molecules are different from those of the dendrimers or the block copolymers. By rationally

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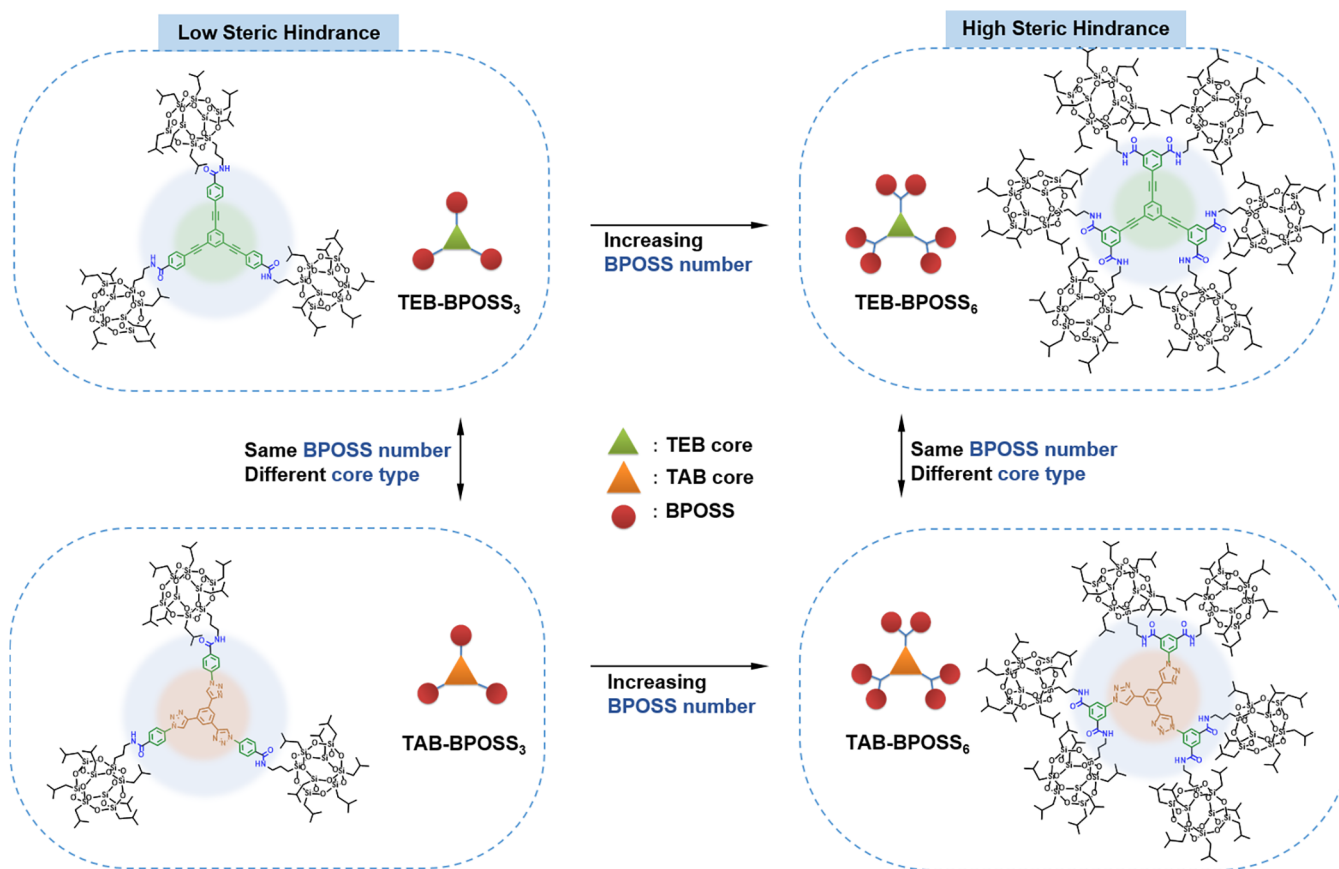


Figure 1. Chemical structures and cartoon illustrations of four giant molecules: **TEB-BPOSS₃**, **TEB-BPOSS₆**, **TAB-BPOSS₃**, and **TAB-BPOSS₆**.

molecular design, various supramolecular nanostructures have been achieved,^{48,50,51} which are largely influenced by the packing constraints of each individual shape as well as competing interactions.⁴⁷ Controlling the packing schemes of these building blocks in a specific combination of composition and topology via a “molecular Lego” approach could result in materials with different structures and properties.^{52–54}

Herein, we propose a strategy to manipulate the packing of nanoscale building blocks via specially designed discotic giant molecules to achieve unconventional supramolecular structures. As a proof-of-concept study, two discotic cores were selected to construct the giant molecules: 1,3,5-triethynylbenzene (TEB, simplified as green triangle in Figure 1) and 1,3,5-triazole benzene (TAB, orange triangles in the cartoons in Figure 1). Connected with flexible chains, both discotic cores are able to form columnar liquid crystal phases with $\pi-\pi$ stacking distances of 0.3–0.4 nm along the column direction. Those cylindrical assemblies have been widely applied into optoelectronic devices and field effect transistors.^{55–57} In this article, three or six relatively rigid and bulky isobutyl POSS (BPOSS) nanoparticles (NPs) (red spheres in the cartoons in Figure 1) are introduced into the periphery of these discotic cores. Since the diameter of BPOSS (1.1–1.2 nm) is much larger than the $\pi-\pi$ stacking distance of the cores (0.3–0.4 nm), therefore, we can directly tune the balance between the steric hindrance of the BPOSS NPs and $\pi-\pi$ interactions of the discotic cores by introducing a different number of rigid BPOSS NPs to the periphery of the discotic cores. When the number of BPOSS is three, the cores may be still able to form columns by rotating the BPOSS NPs around the column. However, when we increase the number of BPOSS to six, the

steric hindrance of the BPOSS NPs at periphery could break the columns down, which may deform into spherical motifs and further self-assemble into spherical supramolecular structures (Figure S1).

Different from our previous works, in which the spherical motifs are formed by aggregation of molecules with cone-shape confirmation whose driving force is the immiscibility between hydrophobic and hydrophilic moieties of the giant molecules,²⁴ in this system, the spherical motifs are formed through breaking down the columns, and the driving forces are the cooperative $\pi-\pi$ interactions and hydrogen bonding competing with the steric hindrance effect. Similar transitions from columnar to spherical phases have been previously observed in dendrimers, which are soft and featured by their generation numbers, molecular tape angle, and solid angle.^{29,31,58} Different from the dendrimers, due to the rigidity of the BPOSS NPs, the steric hindrance could be displayed more directly in the giant molecules. As a matter of fact, there are only limited examples shown that discotic molecules with $\pi-\pi$ interactions could also form F–K spherical packings in dendrimers and giant molecules.^{26,59–62}

2. EXPERIMENTAL SECTION

2.1. Materials. The giant molecules mentioned in the above text were prepared by the Sonogashira reaction or the CuAAC click reaction. More details on the molecular design and synthesis are provided in the SI Appendix, Section 1.2.

2.2. Sample Preparation. Samples with ordered structures were obtained by thermal annealing at 190–230 °C, followed by rapid quenching. Two sample preparation methods, namely drop-casting and microtoming, were used to process TEM samples. For the drop-

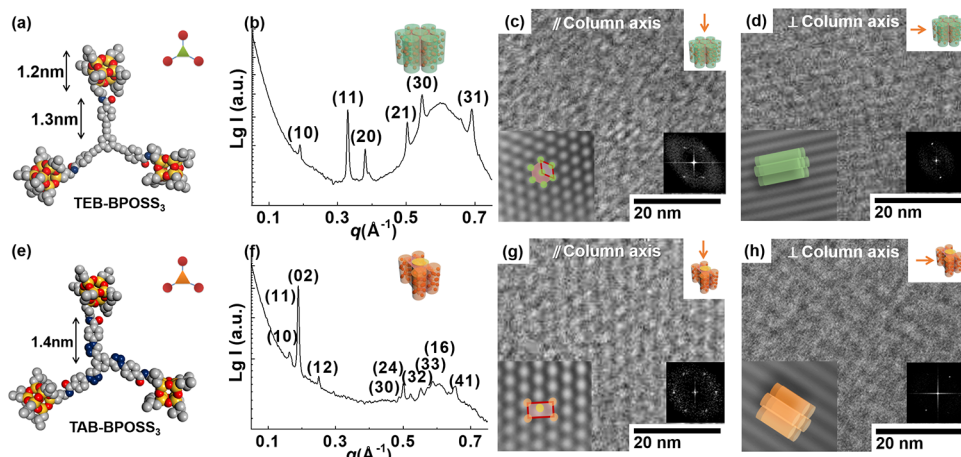


Figure 2. Columnar phases of TEB-BPOSS_3 and TAB-BPOSS_3 . (a) Molecular dimension of TEB-BPOSS_3 . (b) SAXS pattern of TEB-BPOSS_3 . (c,d) Fourier filtered TEM images of TEB-BPOSS_3 taken (c) along and (d) perpendicular to the column axis. Insets: Fourier filtrated TEM images (left bottom) and FFT patterns (right bottom). (e) Molecular dimension of TAB-BPOSS_3 . (f) SAXS pattern of TAB-BPOSS_3 . (g,h) Fourier filtered TEM images of TAB-BPOSS_3 taken (g) along and (h) perpendicular to the column axis. Insets: Fourier filtrated TEM images (left bottom) and FFT patterns (right bottom). The scale bars in the original and Fourier filtered images are the same.

casting method, the sample solution in THF with concentration of 0.5–1.0 mg/mL was dropped onto the carbon-coated copper grids (400 mesh), followed by thermally annealing at corresponding temperature overnight before TEM measurement. For the microtoming method, thin slices of annealed samples for TEM were prepared by utilizing a Leica EM UC7 microtome.

2.3. Characterization. SAXS experiments were performed on a Rigaku MicroMax002+ instrument equipped with a 2D multiwire area detector and a microfocus sealed copper tube. Synchrotron SAXS experiments were conducted at 12-ID-B, C station with X-ray energy of 12 keV at the Advanced Photon Source (APS) of Argonne National Laboratory. Bright field TEM images of the thin-slice samples were collected on a JEOL-1230 TEM with an accelerating voltage of 120 kV and a CCD camera. Fourier filtering of the TEM image was carried out with the reported FFTW implementation method.^{24,63} More details on the characterization are provided in the *SI Appendix*, Section 1.

3. RESULTS AND DISCUSSION

The chemical structures of the four different designed giant molecules are shown in Figure 1. Among them, TEB-BPOSS_3 and TEB-BPOSS_6 possess the same type of core but two different periphery BPOSS numbers (three versus six), and so is the same for TAB-BPOSS_3 and TAB-BPOSS_6 . Their detailed synthetic routes are in the *Supporting Information (Schemes S1–S9)*. Briefly, TEB-BPOSS_3 and TEB-BPOSS_6 are prepared by the Sonogashira coupling of iodo-functionalized BPOSS and alkyne-functionalized 1,3,5-triethynylbenzene cores, while TAB-BPOSS_3 and TAB-BPOSS_6 are synthesized by the copper-catalyzed azide–alkyne cycloaddition (CuAAC) “click” reactions of azo-functionalized BPOSS and alkyne-functionalized 1,3,5-triethynylbenzene cores. The precise chemical structures and monodispersity of these four molecules have been confirmed by ^1H and ^{13}C NMR spectra (Figures S2–S9), matrix assisted laser desorption/ionization-time-of-flight (MALDI-TOF) mass spectra (Figure S14a), and gel permeation chromatography (GPC) results (Figure S14b).

All four samples show excellent thermal stability up to 300 °C (Figure S16). Ordered packing can thus be obtained via thermal annealing at 190 °C–230 °C followed by rapid quenching. Prolonging annealing time (e.g., overnight) does

not change the formed structures. The small angle X-ray scattering (SAXS) technique is used to monitor the formation of ordered structures in the bulk samples. After thermal treatment at 230 °C for 2 min and subsequent quenching, TEB-BPOSS_3 shows a set of well-resolved SAXS peaks (Figure 2b) with a characteristic scattering vector ratio of $1:\sqrt{3}:2$. These peaks can be assigned to a hexagonal columnar lattice, and the projected 2D hexagonal lattice parameters are determined as $a = b = 3.83$ nm and $\gamma = 120^\circ$. The relatively low intensity of the first diffraction peak (10) here is possibly due to the electron density variation within the supramolecular columns, which could also be supported by simplified atomic simulation (Figure S18). Similar phenomena have been attributed to factors such as the presence of a hollow center, variation of the alkyl chain length, introduction of fluorinated chains, and the polyhedral shape of the doubly segregated aliphatic-aromatic supramolecular structure.^{32,64,65}

To further verify this assignment, transmission electron microscopy (TEM) experiments have been carried out for microtomed samples without staining. The scale of the ordered domains are typically several hundreds of nanometers. As shown in Figure 2c, hexagonal columnar packing perpendicular to the column axis can be clearly seen for TEB-BPOSS_3 . The Fourier filtration process is used to enhance the image contrast (see *Supporting Information Section 1.1.2* for details). The intercolumn d -spacing is measured as 3.6 ± 0.1 nm, which is in agreement with that from the SAXS data (3.83 nm). In the TEM image, the darker matrix is attributed to the BPOSS NPs, and the light dots are the stacked TEB cores. Notably, the measured diameter of the light dot is about 2.4 nm, consistent with the diameter of a TEB core (Figure S19). The distance between two light dots is about 1.2 nm, indicating that the BPOSS NPs of adjacent columns may partially interdigitate with each other. TEM images along the [10] direction can also be observed (Figure 2d) with an intercolumn d -spacing of 3.1 ± 0.1 nm, matching well with the (10) d -spacing determined from X-ray data (3.32 nm).

Interestingly, TAB-BPOSS_3 shows a different set of SAXS diffraction peaks with a q -ratio of $\sqrt{3}:\sqrt{4}:\sqrt{7}$ (Figure 2f). These peaks can be assigned as the (10), (11), and (12) planes in a 2D rectangular lattice with lattice parameters of $a = 3.9$

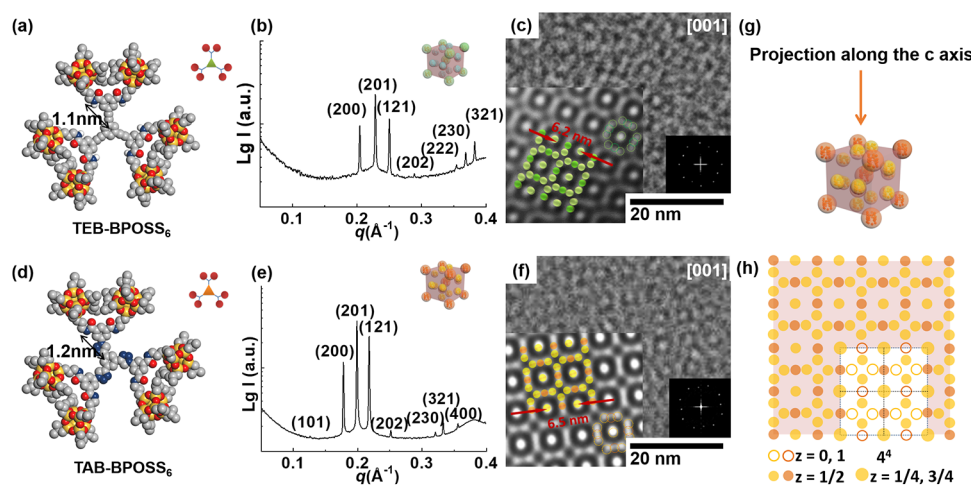


Figure 3. A15 ($Pm\bar{3}n$) phase formation of TEB-BPOSS₆ and TAB-BPOSS₆. (a) Molecular dimension of TEB-BPOSS₆. (b) SAXS pattern and (c) Fourier filtered TEM images of TEB-BPOSS₆. Insets: Fourier filtered TEM images (left bottom) and FFT patterns (right bottom). (d) Molecular dimension of TAB-BPOSS₆. (e) SAXS pattern and (f) Fourier filtered TEM images of TAB-BPOSS₆. Insets: Fourier filtered TEM images (left bottom) and FFT patterns (right bottom). (g) Illustration of the A15 ($Pm\bar{3}n$) structure and (h) the 2D 4⁴ tiling patterns along the [001] direction. The spheres located at sparse layers ($z = 1/4$ and $3/4$) are represented by large filled circles. The spheres at $z = 0$ (small open circles) and $1/2$ (small filled circles) form the dense nets. The scale bars in the original and Fourier filtered images are the same.

nm, $b = 6.64$ nm, and $\gamma = 90^\circ$ and two columns per lattice (for a detailed structure, see Figure S20). This rectangular lattice can also be confirmed by TEM images, as the measured angle of the center column with two neighboring columns deviated from a perfect 60° in a typical hexagonal lattice (Figure 2g). Moreover, lattice parameters determined from this TEM image are $a = 3.9 \pm 0.1$ nm and $b = 6.6 \pm 0.1$ nm, which are in excellent agreement with those obtained from SAXS data. The formation of a rectangular lattice may result from the tilting of the discotic TAB cores within the column, which made the cross section of the columns deviate from circular shape and close to an elliptical shape along the column axis.

Changing the number of BPOSS NPs connected to the triangle core from three to six alters the packing scheme and leads to completely different supramolecular nanostructures. TEB-BPOSS₆ and TAB-BPOSS₆ both show SAXS patterns with three major peaks of q -ratios of $\sqrt{4}:\sqrt{5}:\sqrt{6}$ (Figures 3b and 3e), which are characteristic of a 3D F-K A15 ($Pm\bar{3}n$) phase with spherical motifs.^{6,7} Those diffraction peaks can be indexed as (200), (201), and (121) of the A15 ($Pm\bar{3}n$) lattice. All the other diffraction peaks can also be clearly indexed as in Figures 3b and 3e. The unit cell parameters are determined as $a = b = c = 6.16$ nm, $\alpha = \beta = \gamma = 90^\circ$ for TEB-BPOSS₆ and $a = b = c = 7.06$ nm, $\alpha = \beta = \gamma = 90^\circ$ for TAB-BPOSS₆, respectively. No sharp diffractions can be observed in their WAXD patterns (Figure S21), indicating that BPOSS NPs are not crystallized. Moreover, TEM images taken along the [001] direction exhibit the characteristic 4⁴ tiling patterns for both samples without staining (Figures 3c and 3f–h), confirming the A15 ($Pm\bar{3}n$) structure.²⁴ The light circles refer to the aromatic cores, and the darker area refers to the BPOSS NPs. Spheres of two sizes are observed in the TEM images, suggesting that the A15 ($Pm\bar{3}n$) structures may be the Cr₃Si type.^{66–68} The lattice parameters obtained from TEM images are consistent with those determined from SAXS profiles.

We reason that the changes in the assembled structures from the columns to the A15 ($Pm\bar{3}n$) structure are due to the spatial packing allowed at the periphery of the discotic cores. The discotic cores prefer to stack into columns via $\pi-\pi$ interactions

(Figure 4a). The existence of weak $\pi-\pi$ interactions is supported by the WAXD spectra (Figure S21). It should be

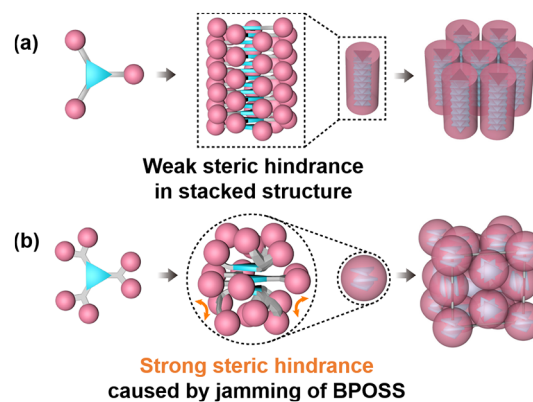


Figure 4. Proposed molecular packing scheme: (a) columnar structure and (b) A15 ($Pm\bar{3}n$) structure. The packing is controlled by the competition between the periphery steric hindrance caused by BPOSS NPs (red spheres) and stacking of the cores (blue triangles).

noted that, in our molecular design, we specifically introduce the amide groups in the linkers, which could generate hydrogen bonding between neighboring molecules and thus further assist the stacking of the discotic cores. The presence of hydrogen bonding can be detected by Fourier transform infrared (FT-IR) spectroscopy (Figure S22). The necessity of this hydrogen bonding in the formation of the structures is illustrated by study on another set of similar molecules with only ester linkages (Schemes S6–S9). The observation that replacing the amide groups by ester linkages completely prevents the formation of ordered structures (Figure S23) highlights the significance of hydrogen bonding for constricting the discotic cores in addition to the weak $\pi-\pi$ interactions.

BPOSS NPs are shape and volume persistent. With only three BPOSS NPs at the periphery of the discotic core (as for TEB-BPOSS₃ and TAB-BPOSS₃), there is enough space for BPOSS NPs to arrange themselves around the core stem by

rotating the molecule along the column direction, so the columnar motifs are retained.³ However, when the number of BPOSS NPs increases to six per core (as for **TEB-BPOSS₆** and **TAB-BPOSS₆**), the NPs at the periphery region are too crowded to maintain the columnar structure. The steric hindrance generated by BPOSS NPs plays a counter role to maintaining the columns. As a result, the columns may prefer to periodically break down along the column axis for releasing the steric hindrance (Figure 4b). In order to balance the extra-volume requirement and the interactions among discotic cores, a bowl-shaped deformation may occur as shown in Figure 4b, leading to formation of spherical motifs, which could further self-assemble into the F-K A15 (*Pm* $\bar{3}$ *n*) phase. A similar mechanism has also been reported in the self-assembly of some dendronized discotic molecules, a different class of molecules, in which their deformation from planar conformation into a bowl shape is induced by the changes in the packing mode of the outer aliphatic region upon temperature increase, leading to the transition from columnar to spherical phases.^{59–62,69,70} We would expect that discotic molecules modified with other structural units with persistent shape and volume (e.g., fullerenes) could also assemble in this way. The measured density values of these two A15 (*Pm* $\bar{3}$ *n*) phases are 1.10 g/cm³ for both **TEB-BPOSS₆** and **TAB-BPOSS₆**. Therefore, in each unit cell, it can be calculated that there are about 24 molecules for **TEB-BPOSS₆** and 30 molecules for **TAB-BPOSS₆**, respectively. Since there are eight motifs in each lattice cell of the A15 (*Pm* $\bar{3}$ *n*) structure, the average numbers of the molecules in each spherical motif are three to four molecules for **TEB-BPOSS₆** and **TAB-BPOSS₆** (see Supporting Information Section 3 for details). This result indicates that in these two samples, the column breaking takes place in every three or four molecules along the imaginary columnar stacking.

4. CONCLUSION

In conclusion, a new strategy performed by symmetric giant molecules has been successfully utilized to construct supramolecular nanostructures with different symmetries. Using a nanoscale discotic core and rigid BPOSS NPs as the prototype models, we show that the balance between the stacking of the cores and the steric hindrance of BPOSS plays a critical role in the formation of self-assembled nanostructures, such as cylinders or spherical Frank–Kasper A15 phases. The rigidity of the BPOSS NPs makes the steric hindrance controlled more directly and simply here. The supramolecular motifs can be tuned from cylindrical to spherical by simply increasing the BPOSS number. This strategy is not limited to BPOSS NPs or the triangle cores. Incorporation of other nanoparticles such as fullerenes, polyoxometalates, and proteins with other joint units can further diversify the packing schemes of such nano building blocks, which might provide insights toward the development of functional materials with specific nanostructures and unexpected properties.

■ ASSOCIATED CONTENT

SI Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acsanm.0c00231>.

Details of the synthesis, chemical structure characterization, additional results, and analysis (PDF)

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Notes

The authors declare no competing financial interest.

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