A Noncrystallization Approach toward Uniform Thylakoids-like 2D “Nano-coins” and Their Grana-like 3D Suprastructures

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Supporting Information

ABSTRACT: Two-dimensional (2D) circular shape nanostructures (e.g., “nano-coins”) are ubiquitously present in thylakoids and grana within chloroplasts of plant cells in nature. The design and fabrication of 2D nano-coins with controlled sizes and thicknesses yet remain challenging tasks. Herein, we present a noncrystallization approach to achieve 2D nano-coins from assemblies of a set of zwitterionic giant surfactants. Distinguished from traditional crystallization approaches where the 2D nanostructures with specific crystallographic symmetries are fabricated, the noncrystallization assembly of giant surfactants results in 2D nano-coins that are derived from the separation of assembled 3D multiple lamellar cylindrical colloids with uniform diameters. The diameters and thicknesses of these nano-coins can be readily tailored by varying the molecular length of giant surfactants’ tails. The formation of 2D nano-coins or 3D cylindrical colloid suprastructures is controlled by tuning the pH value of added selective solvents. This new strategy opens a door for controlling the shape, size, and size distribution of assembled nanostructures with different hierarchies.

INTRODUCTION

Today hierarchically assembled two-dimensional (2D) nano-structures have been extensively studied due to their unique dimensional-dependent properties.1 Most of current techniques for preparing 2D assembled nanostructures involve crystallization processes at certain stages, such as 2D hierarchical nano-sheets formed by crystallization-driven assembly of homopolymers and block copolymers,2 2D nanocrystals of molecular Janus particles1 or peptoid polymers,4 and the generation of dynamic 2D protein crystals from rotational symmetric proteins.5 The overall shapes of these 2D nanostructures are thus dominated by crystallization to form flat Wulff shapes which possess a C2, C3, C4, or C6 rotational symmetry as a result of the crystal unit cell symmetry (Figure 1a),2,4,6 and the size of resultant 2D crystals is rarely controlled, if possible, in a self-seeding process.4 Despite these successes, methods to prepare circular-shaped 2D nanostructures are still largely unexplored, since this type of structures does not contain crystallographically defined lateral surfaces and, thus, requires a noncrystallization approach. In certain assembling conditions, spherical micelles could be deformed and exhibit disk-like quasi 2D nanostructure,8 however, their size, size distribution, and thickness control remain grand challenges. We are devoted to the preparation of precisely controlled circular-shaped 2D nanostructures (like “nano-coins”), which ubiquitously present and play critical roles in nature, such as in chloroplasts of plant cells.9 One chloroplast contains many cylindrical structures (called grana), and each cylinder is formed via stacking of 2D nano-coin thylakoids (Figure 1b), acting to trap the energy from sunlight.10 These unique nano-coin thylakoids and their stacking grana ensure an extremely large area-to-volume ratio for photosynthetic reaction, a high stability, and flexibility in response to dynamically changed environmental conditions.11

In this article, we report a simple approach to achieve well-defined 2D nano-coins via designed zwitterionic giant surfactants. Inspired by 2D crystals based on molecular Janus nanoparticles reported by our group,3 this series of zwitterionic giant surfactants is composed of ionic heads having both positively and negatively charged fullerenes (C60) and a hydrophobic nonpolar polystyrene (PS) tail (abbreviated as NC60−AC60−PSn, Figure 1c), which yet feature noncrystallinity. The molecular design of these zwitterionic giant surfactants captures the essential structural feature of small-molecular zwitterionic surfactants,12 but with amplified molecular sizes.13 In the meantime, pH-responsive groups on the heads of giant surfactants afford the response capability of external conditions. After self-assembling in solution, highly uniform
2D circular shaped nano-coins with controlled lateral sizes and thicknesses are achieved. These 2D nano-coins are able to further stack together to form cylindrical 3D suprastructures at certain conditions. The hierarchically assembled structures resemble the features of thylakoids and grana structures in chloroplasts.

RESULTS AND DISCUSSION

Molecular Design. A series of zwitterionic giant surfactants with different molecular tail lengths of PS were synthesized by a sequential copper-catalyzed azide–alkyne “click” reaction [see Supporting Information (SI), Scheme S1]. Their well-defined chemical structures and high purities were fully characterized (Figures S1–S6). The precursor of NC_{60}−AC_{60}−PS_{6} (compound 7) bears a tert-butoxy carbamate (Boc)-protected C_{60} and a tert-butyl-protected C_{60}. After treating with excessive trifluoroacetic acid (CF_{3}COOH), both Boc and tert-butyl groups were readily hydrolyzed, affording positively charged amino groups with CF_{3}COO− groups as counterions^{14} and neutral carboxylic groups.\[^{15}\] Since carboxylic acid groups can be partially ionized in solution,\[^{16}\] the zwitterionic attributes of giant surfactants are endowed as dissolving in polar solvents where the degree of ionization of carboxylic acid groups can be tuned by the concentration of free hydrogen ions in solution. Such molecular design renders the giant surfactants versatile assembling structures controlled by the pH value of solution.

Formation of Thylakoid-like 2D Nano-coins. Two parameters were focused on in our investigation of solution-assembly behaviors of this series of zwitterionic giant surfactants: an internal structure parameter of the length of PS tail in these surfactants, and an external parameter of the pH value of added aqueous selective solvents. The first experiment was performed by adding HCl aqueous solution (pH = 5, 1.5 mL) into THF (comment solvent) solution of NC_{60}−AC_{60}−PS_{6} (c = 1 mg/mL, 1 mL) at 20 °C (see details in SI). After stirring the mixture at 20 °C for 2 h, air-dried samples were examined by transmission electron microscope (TEM). Uniform individual 2D nano-coins with a diameter around 250 nm were observed (Figure 2a). Based on the results obtained from atom force microscopy (AFM), the thicknesses of these nano-coins were measured to be 15.3 ± 0.2 nm (Figure 2b and c).

We have confirmed that individual nano-coins can also be observed by adding HCl aqueous solution with their pH values ranging between 3 and 5 into NC_{60}−AC_{60}−PS_{6}/THF solution (Figures S7 and S8). We also noted that the 2D nano-coins structures were stable over a period of three months. When the length of PS tail was decreased to 62 repeating units, an individual nano-coin with a diameter of 250 nm was also observed (Figure 2d), while the thickness of the resulting individual nano-coin is decreased to 12.7 ± 0.3 nm, as measured by AFM (Figure 2e and f). When the length of PS tail was increased to 120 repeating units, nano-coins with a thickness of 17.5 ± 0.5 nm were found (Figure S9) but, however, have a relatively wide distribution in their diameters (Figure S10). If considering that the diameter of ionized C_{60} is around 2 nm\[^{14,16}\] and the radii of gyration (R_g) of PS are 2.18 and 2.57 nm for PS_{62} and PS_{6} (see SI for calculations), respectively, the resulting individual thicknesses of nano-coins can be qualitatively judged that they consist of a single layer of zwitterionic giant surfactant heads at both of the top and bottom, and two layers of the PS tails are sandwiched in between as illustrated in Figure 2g and h. Furthermore, wide-angle X-ray diffraction (WAXS) and electronic diffraction (ED) results of 2D nano-coins confirm their noncrystalline feature within the nano-coins (Figure S11).

To understand the formation pathway and mechanism of the nano-coins, dynamic light scattering (DLS) experiments were carried out to investigate their solution assembly behaviors. Typical CONTIN analyses of DLS measurements for these three samples indicate that the formation of assembled structures have an average hydrodynamic radius (R_h) of 350 nm with minor angular dependence (Figure S12a for NC_{60}−AC_{60}−PS_{6} as an
example), suggesting the presence of close to isotropic aggregates in the solution. Their $R_c$ values are much larger than the size of individual 2D nano-coin, further revealing that the stacked rather than well-dispersed nano-coins present in the solution (Figure S12a for NC$_{60}$^\textsuperscript{−}−AC$_{60}$^\textsuperscript{−}−PS$_{62}$ as an example). The detailed internal structure of these stacking of 2D nano-coin structures for NC$_{60}$^\textsuperscript{−}−AC$_{60}$^\textsuperscript{−}−PS$_{62}$ is revealed by cryo-TEM imaging without staining. As shown in Figure 3a, cylinder-like colloids having multiple layers are observed. The building blocks are layered structures with identical thickness and diameter (lateral size). Since the electron density of C$_{60}$ is higher than that of PS, the dark layers are attributed to the aggregation of the zwitterionic C$_{60}$ heads (4 nm). The combination of the DLS and cryo-TEM results verifies the formation of multiple layered cylindrical colloids rather than individual nano-coins in solution. Fourier transform infrared (FTIR) results on the freeze-dried powder samples of NC$_{60}$^\textsuperscript{−}−AC$_{60}$^\textsuperscript{−}−PS$_{62}$ reveal around 28% of carboxylic groups are deprotonated (Figure S13), indicating that surfaces of the nano-coins are overall positively charged, which is further confirmed by zeta potential result of sample (for NC$_{60}$^\textsuperscript{−}−AC$_{60}$^\textsuperscript{−}−PS$_{62}$, +65.3 mV) in solution. The static charge repulsion between the C$_{60}$ layers may lead to a relatively loose layer packing. All of these experimental results suggest that the nano-coins are derived from the separation of cylindrical colloids containing multiple layered structures, reminiscent of the falling of a cylindrical column of well-stacking coins. A direct evidence of this assessment is that we can occasionally observe in real space the existence of partially fallen nano-coins with one stacked on another under cryo-TEM for steered NC$_{60}$^\textsuperscript{−}−AC$_{60}$^\textsuperscript{−}−PS$_{62}$ solution (Figure 3c). Furthermore, we can also observe under bright-field TEM for the same samples; in many cases, uniform nano-coins are lying down and partially stacked one by one, similar to the case after “circular Domino in falling” as shown in Figure 2d. The similar observation for NC$_{60}$^\textsuperscript{−}−AC$_{60}$^\textsuperscript{−}−PS$_{62}$ is also clearly presented in Figure 2e. The separation of individual “coins” is attributed to the charge repulsion between layer surfaces of ionized C$_{60}$, yet further revealing that the two zwitterionic C$_{60}$ heads in one giant surfactant are located in the same C$_{60}$ head layer of the “coin” due to the PS tail chemical linking location to the C$_{60}$ head (Figure 2h). Otherwise, if there is a connection between two C$_{60}$ head layers, this separation of the individual coin could not be possible. The formation pathway is thus proposed in Figure 3f. The nano-coins are generated via nanophase separation between C$_{60}$ and PS tails inside the cylindrical colloids and thus possess identical diameters. Due to the loose packing between two C$_{60}$ layers, they will fall down and form a “circular Domino in falling” after solvent evaporating as observed in Figure 3d and e.

**Figure 2.** Morphological characterization of 2D nano-coins. (a) Bright-field TEM images of NC$_{60}$^\textsuperscript{−}−AC$_{60}$^\textsuperscript{−}−PS$_{62}$ prepared by adding aqueous HCl solution (pH = 5) into giant surfactants in THF solution. (b) AFM image of NC$_{60}$^\textsuperscript{−}−AC$_{60}$^\textsuperscript{−}−PS$_{62}$ in the height mode. The upper-left inset is the amplitude image in selective area. (c) The height profiles of 2D nano-coins in panel b. (d) Bright-field TEM images of NC$_{60}$^\textsuperscript{−}−AC$_{60}$^\textsuperscript{−}−PS$_{62}$ prepared by adding aqueous HCl solution (pH = 5) into giant surfactants in THF solution. (e) AFM image of NC$_{60}$^\textsuperscript{−}−AC$_{60}$^\textsuperscript{−}−PS$_{62}$ in the height mode. The upper-left inset is the amplitude image in upper-right selective area. (f) Height profiles of 2D nano-coins in panel e. (g) Schematic representation and (h) molecular packing of a 2D nano-coin formed by an assembly of zwitterionic giant surfactants.
It should be pointed out that, once the cylindrical colloids are formed in this condition, further adding lower pH values of HCl aqueous solution will not alter the final colloidal morphology. No separation of nano-coins was observed after the samples were dried and prepared for TEM observations.

With increasing the length of the PS-tails to $n = 86$, the cylindrical colloids with stacked layers also possess a diameter of around 250 nm as identified by TEM (Figure 4b) and SEM (Figure 4b, inset) observations. In theory, when the size of PS tail is much bigger than that of the zwitterionic C$_{60}$ heads, the assembly behavior of this molecule will be dominated by PS tails. In this case, the surface free energy of colloids dominates the assembling process, and the overall shape becomes spherical. Indeed we found that, when the PS tail with 120 repeating units was used, spherical colloids with partial layer stacks near the center of the colloids were observed, yet they possess a diameter of 220 nm (Figure 4c). It can be expected that if the “circular Domino in falling” collapsing of the cylinder colloids takes place as the overall positive surface charge of the C$_{60}$ layer increases (decreasing the pH value of added HCl aqueous solution), uniform 2D nano-coins would be produced. If the colloid shape deviates from cylindrical (such as becoming spherical), the 2D nano-coins with a broad distribution of their lateral sizes-diameters can be expected.

Figure 4d–f shows bright-field TEM images for three cylindrical colloids with a high-magnification in the selective area of Figure 4a–c. Their corresponding fast Fourier transform (FFT) patterns are also inserted in these figures. When we compare the nano-coins stacked structures formed in these three samples, it is found that the thicknesses of the darker C$_{60}$ layers keep constant at around 4.0 nm, consistent with a double size of diameters of the close-packed ionized C$_{60}$ heads. Those of greyer PS layers increase from 8.6 nm, to 11.0 to 12.5 nm, with an increasing of PS tail length from $n = 62$, to $n = 86$, and further to $n = 120$, respectively.

The colloid growth process can be monitored by TEM at different water contents (Figure S14). As neutral water was added into the NC$_{60}$–AC$_{60}$–PS$_{62}$/THF solution, hydrophobic PS tails tend to aggregate and stimulate nanophase separation between ionized C$_{60}$ heads and nonpolar PS tails due to the incompatibility between two components. The growth direction along the layer normal is governed by the aggregation of the PS tails and zwitterionic C$_{60}$ heads which have been neutralized by the counterions in solution. The lateral growth perpendicular to the layer normal is attributed to accumulative addition of the zwitterionic giant surfactants into assemblies. When the PS tails are overcrowded, they could be squeezed out from the PS layers and eventually cover the edges of the C$_{60}$ head ionic layers to prevent their further lateral growth. The overall shape of the colloids is governed by minimizing overall surface free energy.

The squeezed PS tails can be described by the concept of reduced tethering densities ($\tilde{\sigma}$) of PS tethered on a flat surface. It is defined by

$$\tilde{\sigma} = \frac{\sigma}{R_g^2}$$  \hspace{1cm} (1)

where $R_g$ is the radius of gyration of PS chain, and $\sigma$ is the tethering density, defined as the reciprocal average covering area of each chain. It has been experimentally proved that the tethered chains start to be squeezed by their neighboring chains at a critical $\tilde{\sigma}^* \sim 3.7$. The $\tilde{\sigma}$ values for our three samples have been calculated to be 4.1, 5.8, and 8.1 (see calculations in SI),

![Figure 3. Formation pathway and mechanism of the 2D nano-coins.](image)

(a) Cryo-TEM image of NC$_{60}$–AC$_{60}$–PS$_{62}$ in THF/aqueous HCl (pH = 5) solution, where the nano-coins are stacking together. (b) High-magnification cryo-TEM image for the selective area shown in panel a. (c) Cryo-TEM image of steered NC$_{60}$–AC$_{60}$–PS$_{62}$ solution, where the partially separated and stacked nano-coins are presented. The inset is the cartoon illustrating the structures in the cryo-TEM image. Bright-field TEM images of (d) NC$_{60}$–AC$_{60}$–PS$_{62}$ and (e) NC$_{60}$–AC$_{60}$–PS$_{62}$ in THF/aqueous HCl (pH = 5) solution, where the 2D nano-coins are stacking one by one, resembling a collapse of a stack of ‘coins’. (f) Schematic illustration of the discrete nano-coins when samples are dried out.

Figure 3. Formation pathway and mechanism of the 2D nano-coins.
respectively, indicating that the PS tails are overcrowded. If the layer lateral dimension is infinitely large, the squeezed PS tails must be stretched orthogonal to the layer (Figure 5, in bulk state), leading to an increase of the layer spacing. However, if the lateral layer dimension starts growing at a limited size such as in the present case, the PS tails could be squeezed out at the edges of the layers (Figure 5, in solution state). The PS tail coverage on the periphery of C60 ionic layer edges can be directly observed in bright-field TEM images (arrows in Figure 4). Specifically, a more clear observation is available from high-magnification TEM images (Figure S1S), where a PS layer with thickness of ca. 12.5 and 11.9 nm is covered in the front of the C60 ionic layer in the case of NC60−AC60−PS62 and NC60−AC60−PS86, respectively.

To quantitatively understand the overcrowding and squeezing out of the PS tails, we compare the d-spacing of the layers in the bright-field TEM images (arrows in Figure 4). Specifically, a more clear observation is available from high magnification TEM images (Figure S1S), where a PS layer with thickness of ca. 12.5 and 11.9 nm is covered in the front of the C60 ionic layer in the case of NC60−AC60−PS62 and NC60−AC60−PS86, respectively.

Figure 4. Formation of cylindrical colloids of stacked 2D nano-coins and the comparison of their lamellae thicknesses with those formed in bulk states. Bright-field TEM images of (a) NC60−AC60−PS62, (b) NC60−AC60−PS86, and (c) NC60−AC60−PS120, prepared by adding neutral water (pH = 7) into their THF solutions, respectively. Insets are the corresponding SEM images. Panels d, e, and f are high-magnification TEM images of the selective area in panels a, b, and c. The insets are the FFT patterns of the TEM images. SAXS patterns of (g) NC60−AC60−PS62, (h) NC60−AC60−PS86 and (i) NC60−AC60−PS120 in bulk states, prepared by thermal annealing at 130 °C for 12 h.

Figure 5. Stretching and squeezing of PS in the bulk and solution state. In bulk state, PS chains are stretched orthogonal to the C60 layer, where a radius (r) is assigned and the thickness (d1) can be obtained from SAXS results. In solution state (cylindrical colloids), the PS chains are squeezed out at C60 layer edges. The thickness (d2) can be obtained from TEM images in Figure 4. After squeezing, the radius of column is r + t, where t is the PS layer thickness surrounding the column and can be measured from TEM image in Figure S15.
cylindrical colloids with those formed in the solid state of these zwitterionic giant surfactants. After thermally annealed at 130 °C for 12 h in the bulk, well-ordered, layered structures were observed for all three samples of NC60−AC60−PS62, NC60−AC60−PS86, and NC60−AC60−PS120, evidenced by the q-ratios of 1:2.3:4.5 in the small-angle X-ray scattering (SAXS) patterns (Figure S13), indicating that the multiple layered structures are formed by the nanophase separation of the C60 ionized heads and PS tails. Since the PS tails in the bulk cannot be squeezed out at the layer edges, the d-spacings of the layers formed in the bulk must be larger than those in cylindrical colloids. This is indeed the case as experimentally observed: 15.7 nm vs 12.6 nm for NC60−AC60−PS62, 18.5 nm vs 15.0 nm for NC60−AC60−PS86, and 21.6 nm vs 16.5 nm for NC60−AC60−PS120, respectively, as summarized in Table 1. Therefore, the PS tails in the bulk state are expected to be more stretched than those in cylindrical colloids. The stretching ratio (S) of PS tails in the cylindrical colloids and the bulk state can be quantitatively characterized by:

\[
S = \frac{L}{2R_g}
\]

where \(L\) is the half layer thickness of PS domain. The S values are close to unity in cylindrical colloids (Si values in Table 1) for these three samples, while they range from 1.35 to 1.45 in the bulk state (Sf values in Table 1). These results indicate that the PS tails are stretched in the bulk state but not so in the cylindrical colloids. To minimize free energy, the PS tails in the cylindrical colloids tend to be released and squeezed on to free layer edges. This type of tendency is accumulative.

In view of a constant PS density in both cases, the volume of PS tails (\(V_p\)) that are squeezed out at the layer edges can be calculated by,

\[
V_p = \pi r^2 (d_1 - d_2)
\]

where \(r\) is a radius of an assumed column in the bulk and \(d_1\) and \(d_2\) are periodic lamellar d-spacing in the bulk and cylindrical colloids, where they include one PS domain and two layers of C60 (Figure S16). Note that the C60 layer thickness in the bulk state is also ca. 4.0 nm (Figure S16), identical to that in the cylindrical colloids. The volume of PS layer covered on the lateral growth surface of the C60 ionic layers can be calculated by

\[
V_p = \pi (r + t)^2 - \pi r^2 \times d_2
\]

where \(t\) is the thickness of PS layer along the radial direction covering on the lateral growth surface of the C60 ionic layers (Figure S16).

When one has \(V_p = V_p\), indicating the lateral growth surface of C60 ionic layers are completely covered by the PS tails, a radius of cylindrical colloids \((r + t)\) can be calculated to be 120.0 nm for NC60−AC60−PS62 and 119.5 nm for NC60−AC60−PS86, or their diameters are 240.0 and 239.0 nm, respectively (see calculation in the SI). They are close to our experimental observation of 250 nm as shown in Figure 4a–b.

Assembling Behaviors under Other pH Values. Using HCl aqueous solution (\(pH \leq 2\)) as selective solvent, regular micellar behaviors of giant surfactants in solution were exhibited. For example, by adding HCl aqueous solution (\(pH = 1\)) into the THF solution of the giant surfactants (\(c = 1\) mg/mL) at 20 °C, the ionization of carboxylic groups in the head are completely suppressed (Figure S13), and only positively charged C60 heads are functioning with hydrophobic PS tails in the assembly process. We thus observe the normal micellar formations, and they change from spheres, to cylinders, further to vesicles with increasing the length of PS tails from NC60−AC60−PS62 to NC60−AC60−PS86 and further to NC60−AC60−PS120 (Figure S17), consistent with the solution assembly behaviors of our previously reported giant surfactants.13b,15b

**CONCLUSIONS**

In summary, we report a noncrystallization self-assembly approach to achieve unique round-shape 2D nano-coins with controlled sizes and thicknesses from zwitterionic giant surfactants. Differentiating from the crystallization-driven approaches where delicate control of crystallization conditions is essential and each individual single crystal 2D nanostructure grows independently, the self-assembly process of zwitterionic giant surfactant is readily directed by the strong nanophase separation between ionized C60 heads and nonpolar PS tails to generate the 3D cylindrical colloids with internal disk-stacked structures. The noncrystallization anisotropic growth lead to the round shape of cylindrical colloids, whose sizes is controlled by the full coverage of overcrowding and squeezing out PS layer on the periphery. These 3D cylindrical colloids ensure the uniform diameter of nano-coins. More importantly, the as-formed 3D cylindrical colloids can either be stabilized as suprastructure with controllable morphologies, or separated into 2D nano-coins by simply tuning pH value of the added selective solvents. This hierarchical assembly pathway resembles 2D nano-coin thylakoids stacking together to form 3D cylindrical granules in chloroplasts. This strategy is easily practicable and expected to be readily applicable to many other noncrystalline polymer systems. In addition to using templates and controlling kinetics during assembling process (e.g., living self-assembling process), this work affords us a new methodology to control the shape, size, and size distribution of assemblies in solution.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.7b01275.

Experimental details, additional results and discussion (PDF)