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A Block Supramolecular Polymer and Its Kinetically Enhanced Stability

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ABSTRACT: Biomolecular systems serve as an inspiration for the creation of multicomponent synthetic supramolecular systems that can be utilized to develop functional materials with complexity. However, supramolecular systems rapidly reach an equilibrium state through dynamic and reversible non-covalent bonds, resulting in a disorganized mixture of components rather than a system in which individual components function cooperatively and/or independently. Thus, efficient synthetic strategies and characterization methods for intricate multicomponent supramolecular assemblies need to be developed. Herein, we report the synthesis of porphyrin-based supramolecular polymers (SPs) in which two distinct block segments consisting of different metal porphyrins are connected: i.e., block supramolecular polymers (BSPs). BSPs with a controlled length and narrow polydispersity were achieved through seeded-growth by a solvent mixing protocol. Interestingly, the block structure permitted the SP as an inner block to coexist with a reagent that was otherwise incompatible with the SP alone. We infer that the inner SP block is compartmentalized in the block structure and endowed with the kinetic stability. Molecular simulations revealed that monomer exchange occurs from the termini of the SP, which corroborated the enhanced stability of the BSP. These results are expected to pave the way for the design of more complex multicomponent supramolecular systems.

INTRODUCTION

In the light of the substantial contribution of living polymerization to polymer science^{1,2}, the recent development of its supramolecular counterpart, i.e., living supramolecular polymerization^{3,4}, has inspired us to expand the scope of molecular self-assembly in the fields of materials science, nanotechnology, and biotechnology⁵⁻ ⁷. In this regard, the synthesis of block supramolecular polymers (BSPs), which offers unprecedented structural complexity and sophisticated functionality, has attracted considerable attention despite it being significantly challenging⁸.

Based on the previous studies⁹⁻²⁵, we infer the prerequisites of living supramolecular polymerization and the issues associated with the synthesis of BSPs. First, the mechanism²⁶⁻²⁹ nucleation-elongation should be operative so that supramolecular polymerization can be processed in a manner analogous to chain growth polymerization. In general, such supramolecular polymers (SPs) have high degrees of internal order (or crystallinity); accordingly, the synthesis of BSPs appears as difficult as welding two single crystals of different molecules. To this end, it is necessary to consider interfacial design between the two blocks, including intermolecular interactions and lattice matching³⁰. Second, a metastable state has to be involved such that an otherwise spontaneous nucleation

process is kinetically suppressed. Formation of a competing aggregate^{9,10} and intramonomer hydrogenbonding^{11-14,20,21} have been demonstrated to be effective for this purpose. However, it is still difficult to manage the pathway complexity because the delicate balance between the stabilities of the metastable species and SPs gets readily disrupted if the monomer structure is altered^{10,13,31}. Given the abovementioned considerations, monomer design of the second block in a BSP is inevitably restricted by the monomer structure of the first block.

One-dimensional (1D) block nanostructures have been reported to date^{8,32-40}, and in particular, living crystallization-driven self-assembly (CDSA) has been demonstrated to be a powerful method to obtain such architectures via self-assembly³⁴⁻³⁷. Nevertheless, 1D block nanostructures with a unimolecular width (i.e., BSPs) are quite rare^{8,10,15,17}. Previously, van der Gucht, Otto, and coworkers¹⁵ synthesized a BSP of peptide-based macrocyclic monomers bearing benzyl and cyclohexylmethyl side chains. We have also achieved a BSP of porphyrin monomers bearing methoxy (2_{Zn}Me) and hexyloxy $(2_{Zn}Hex)$ side chains (Figure 1)¹⁰. These examples imply that monomer structures in a BSP have to be very similar, which corroborates the aforementioned assertion. In fact, regarding our previous porphyrin systems, even a BSP consisting of different metal porphyrin cores (e.g., 2ZnMe and 2_{Cu}Me) could not be obtained (S. Ogi, M. Takeuchi, K. Sugiyasu, unpublished results). Although Palmans, Meijer, and co-workers recently reported characterization of BSPs under thermodynamic control⁸, the functions and properties of BSPs arising from dissimilar SP blocks have thus far been virtually unexplored.

Herein, we assessed the applicability of the living CDSA method to the synthesis of unimolecular BSPs. In this protocol, seeded-growth is performed by mixing good and poor solvents for monomers. Manners and coworkers extended their living CDSA method to the supramolecular polymerization of platinum complexes and succeeded in controlling the length of SPs^{16,17}. Furthermore, they succeeded in creating a unique scarflike nanostructure consisting of the platinum complexes bearing polyethylene glycol side chains with different lengths (i.e., $(C_2H_4O)_n$, where n = 12 or $16)^{17}$. For the synthesis of BSPs, de Greef, Meijer, and co-worker41 proposed a stepwise assembly of multicomponent structures by using good and poor solvents based on their in-depth mechanistic study; however, this strategy has not been attempted as yet.

We succeeded in connecting the SPs consisting of distinct porphyrin cores in B–A–B type triblock copolymer structures through the solvent mixing protocol. Interestingly, it was found that the stability of block A could be kinetically enhanced by the presence of block B, which illustrates a collective property of BSPs dictated by the block sequence. This study represents an important step toward the synthesis of advanced multicomponent SPs that have been increasingly attracting attention in recent years⁴²⁻⁴⁷.

RESULTS AND DISCUSSION



Figure 1. Structure of porphyrin-based monomers.

Molecular design. As mentioned above, BSPs of different metal porphyrin monomers could not be obtained in our previous study (e.g., by using 2ZnMe and 2CuMe) because these monomers showed different selfassembling behavior (Figures S5 and S6 for 2ZnMe and 2_{Cu}Me, respectively). We reasoned that these porphyrin cores are distorted differently and adopt different stacking modes⁴⁸. To address this issue, we increased the number of hydrogen bonding sites so that the junction between the different metal porphyrin cores could be matched and stabilized. The synthesis and

characterization of the new monomers, namely, $4z_n$ and $4c_u$, (Figure 1), are described in the Supporting Information.

Supramolecular polymerization. As discussed later, seeded-growth in living CDSA is typically performed in a mixed solvent system. In this study, a mixture of methylcyclohexane (MCH): a poor solvent that induces self-assembly of porphyrin monomers, and toluene: a good solvent in which porphyrin monomers are molecularly dissolved, was used. Figure 2 illustrates the supramolecular polymerization of the zinc(II)-porphyrin monomer ($4z_n$) in mixed solvents.



Figure 2. Self-assembling behavior of 4_{Zn} in MCH/toluene mixed solvents. (a) Absorption spectra of 4_{Zn} in MCH/toluene mixed solvents: $[4_{Zn}] = 10 \ \mu$ M. (b) AFM image of supramolecular polymer of 4_{Zn} (i.e., SP4_{Zn}) prepared in MCH/toluene (2:1) mixed solvent: scale bar, 400 nm. (c) Plot of absorbance at 398 nm as a function of MCH proportion.

On increasing the proportion of MCH, the Soret band blue-shifted (Figure 2a), which suggested that the porphyrin chromophores were self-assembled in a faceto-face stacking manner, i.e., H aggregate mode^{9,10}. Atomic force microscopy (AFM) visualized the 1D supramolecular polymer (Figure 2b). The plot of absorbance at 398 nm as a function of the proportion of MCH showed a critical MCH/toluene ratio as ~40% of MCH, below which complete disassembly occurs (Figure 2C). This result implied that supramolecular polymerization of 4zn proceeded via the nucleationelongation mechanism^{41,49,50}. In fact, temperaturedependent absorption spectral changes (cooling process) accompanied a critical elongation temperature (T_e) , a characteristic of the nucleation-elongation process (Figure 3a,b)⁵¹⁻⁵³. We found a linear relationship between the $T_{\rm e}$ values and the proportion of MCH (Figure 3c).

Figure 3b shows the temperature-dependent degree of supramolecular polymerization obtained for solutions of $4z_n$; the results obtained for 67% MCH solution are

shown as typical cooling curves. The elongation regime, i.e., the curve below $T_{\rm e}$, could be fitted by the model developed by van der Shoot and Meijer⁵¹. The elongation enthalpy ($H_{\rm e}$) determined from curve fitting was –108 kJ mol⁻¹ for the curve of [4zn] = 10 µM. The standard enthalpy (ΔH°) and entropy (ΔS°) for the elongation process were determined from a van't Hoff plot to be –119 kJ mol⁻¹ and –271 J mol⁻¹ K⁻¹, respectively (Figure S7). Thus, the Gibbs free energy (ΔG°) at 298 K was calculated to be –38 kJ mol⁻¹. Likewise, 1D supramolecular polymerization of 4_{Cu} was assessed (Figures S9 and S10): ΔH° = –122 kJ mol⁻¹, ΔS° = –284 J mol⁻¹ K⁻¹, and ΔG° (at 298 K) = –37 kJ mol⁻¹.



Figure 3. Supramolecular polymerization of $4z_n$ in MCH/toluene (2:1). (a) Temperature-dependent absorption spectral changes of $4z_n$ in MCH/toluene (2:1): $[4z_n] = 10 \mu$ M. (b) Changes in the degrees of supramolecular polymerization (α) of $4z_n$ in MCH/toluene (2:1) as a function of temperature. A red solid line is the fitting curve for $[4z_n] = 10 \mu$ M, obtained from cooperative model. (c) Plot of T_e values as a function of MCH proportion.

As demonstrated above, the new porphyrin monomers ($4z_n$ and 4_{Cu}) have a strong propensity for supramolecular polymerization because of the increased number of hydrogen-bonding sites. Thus, it is anticipated that block junction between dissimilar SPs could be stabilized to produce BSPs. However, unlike our previous system of $2z_n Me^{9.10}$, the supramolecular polymerization of these new monomers does not involve a metastable state, which eliminated the possibility of kinetic control over supramolecular polymerization (see the prerequisites discussed in the Introduction). As such, a dilemma was inherent in this monomer design, which led us to investigate whether the solvent mixing protocol such as the living CDSA was applicable to the synthesis of unimolecular BSPs^{16,17}.

Living supramolecular polymerization. Figure 4a depicts the scheme of living supramolecular

polymerization performed in this study. Briefly, solutions of the seeds of a SP (in MCH) and a monomer (in toluene) were mixed to initiate the seeded-growth. The plots of the temperatures at which the degree of aggregation reached 97% against the proportion of MCH suggested that $_{4Zn}$ is fully dissolved in toluene but self-assembles in 67% MCH at 298 K ($[_{4Zn}] = 10 \mu$ M, Figure S11). Therefore, provided a sufficiently high monomer concentration, this plot assures that a high degree of aggregation (>95%) can be retained when the solutions of the seeds (in MCH) and the monomer (in toluene) are mixed in a ratio of 2 to 1 at 298 K (i.e., 67% MCH); this mixing ratio is used for the experiments that follow.

The scheme shown in Figure 4a requires seeds of a SP. By sonication (90 W at 2 °C for 3 h), SP4zn fragmented into short pieces with the number average length (L_n) and weight average length (L_w) of 86 and 102 nm, respectively (Figure 4b. Figures S12a-c). The absorption spectrum of these SP4zn seeds was identical to that of SP4zn, indicating that the H aggregate mode was preserved (Figure S12d).

The MCH solution of SP_{4zn} seeds was placed carefully on the top of the toluene solution of monomeric 4zn in a cuvette, which was then shaken for a few seconds at room temperature (Figure S13). The color of the mixed solution instantaneously changed to that of the H aggregate. The kinetics of the seeded-growth was too fast to be monitored even with the stopped-flow setup. The "depolymerization overshoot"⁴¹ of the seed—the temporal depolymerization caused by inefficient mixing of the good and poor solvents—was not observed in both the manual mixing experiment and the stopped-flow experiment (Figure S14).

Figures 4c and d show the AFM images of SP4zn seeded-growth obtained after with different seed/monomer ratios (see also Figure S15). The L_n and L_w values increased in agreement with the feed ratio between the seed to the monomeric 4zn (for 1:1 and 1:5 seed-tomonomer ratios, $L_n = 177$ and 422 nm, $L_w = 214$ and 523 nm, respectively). Thus, the polydispersity indices (PDI, $L_{\rm w}/L_{\rm n}$) were as low as 1.21–1.24, indicative of controlled growth (Figure 4e). The controlled length with narrow polydispersity was retained at least for a week after the sample preparation, suggesting slow dynamic nature of our SPs. It should be noted that such a low PDI value could not be achieved without using the seed, i.e., if a toluene solution of 4zn was mixed with pure MCH in the scheme shown in Figure 4a (PDI = 1.49, Figure S16). Small ill-defined aggregates, which were most probably formed without the aid of the seed, were also observed particularly for the sample prepared at the 1:5 seed-tomonomer ratio (Figure 4d). We assert that the contribution of these aggregates to the following experiments is negligible because the absorption spectrum of the thus-obtained SP4Zn was identical to that of the SP_{4zn} prepared through heating/cooling process under thermodynamic control (Figure 3a, see Figure S17 for detail). It is worth noting that the mixing processes (shaking, stirring, injection, etc.) affected the PDI values, and it was necessary to optimize them as well;



Figure 4. Living supramolecular polymerization and synthesis of block supramolecular polymers. (a) Schematic representation of living supramolecular polymerization performed via the CDSA method. (b) AFM image of seed of SP_{4zn} scale bar, 400 nm. (c,d) AFM images of SP_{4zn} prepared according to the scheme shown in (a): scale bar, 400 nm. The ratios between the seed of SP_{4zn} and monomeric 4_{Zn} are 1:1 (c) and 1:5 (d). (e) Cumulative histograms of the length distributions of (i) the seed of SP_{4zn} and SP_{4zn} prepared with the seed to monomeric 4_{Zn} ratios of (ii) 1:1 and (iii) 1:5. (f,g) AFM images of SP_{4zn} and monomeric 4_{Cu} were 1:1 (f) and 1:5 (g). (h) Cumulative histograms of the length distributions of (i) the seed of SP_{4zn} and SP_{4zn} and SP_{4zn} of the scheme shown in a, scale bar = 400 nm. The ratios of the seed of SP_{4zn} and monomeric 4_{Cu} were 1:1 (f) and 1:5 (g). (h) Cumulative histograms of the length distributions of (i) the seed of SP_{4zn} between 1:1 (f) and 1:5 (g). (h) Cumulative histograms of the length distributions of (ii) the seed of SP_{4zn} and SP_{4zn} between the seed to monomeric 4_{Cu} ratios of (ii) 1:1 and (iii) 1:5.

specifically, rapid and homogeneous mixing of the two solutions achieved lower PDI values. Further studies including the use of a variety of monomers with different structures and the combination of good/poor solvents are currently underway.

In the last decade, Manners and co-workers³⁴⁻³⁷ have developed the living CDSA method and fabricated a variety of block nanostructures consisting of block copolymers. Fukushima, Aida, and co-workers³⁸ synthesized a linear heterojunction nanotubular object composed of small molecular entities (hexabenzocoronene derivatives). These seminal studies have demonstrated the unprecedented potential applications of block nanostructures. Similarly, in the scheme shown in Figure 4a, other monomers such as 4Cu in its toluene solution could be used instead of 4zn. The increased L_n and low PDI values confirmed the controlled seeded-growth, thus suggesting the formation of a block supramolecular polymer, BSP(4cu-b-4zn-b-4cu) (for 1:1 and 1:5 seed(SP_{4zn})-to-monomer(4_{Cu}) ratios, $L_n = 18_3$ and 4_{53} nm, PDIs = 1.21 and 1.21, respectively: Figure 4f,g,h and Figure S18). Interestingly, fluorescence of SP4zn was

quenched by block copolymerization with SP_{4cu}, which indicates electronic communication such as excitation energy transfer between the two blocks (Figure S19)^{38,56}. The obtained BSP consisting of monomers based on distinct cores (not side chains^{10,15,17}) represents a new type of multicomponent SPs^{8,42-47}.

Kinetic stability of the block supramolecular polymers. The reversible nature of non-covalent bonds endows SPs with unique dynamic properties such as stimuli responsiveness and self-healing ability^{5-7,54} However, it also leads to a drawback that SPs are inherently unstable, hindering their further applications to a certain extent. For example, SP_{4zn} was readily depolymerized in the presence of Lewis bases that coordinated to the zinc atom and prevented the porphyrin plane from forming π -stacks^{55,56}. Upon the addition of 4-dimethylaminopyridine (DMAP), the H aggregate absorption band of SP_{4zn} disappeared (398 nm) and a new band corresponding to DMAP-coordinated zinc(II)-porphyrin could be observed (430 nm) (Figure 5a). The Gibbs free energy for the coordination of DMAP to zinc(II)-porphyrin (1zn) in the MCH/toluene (2/1)

mixture was determined to be -29 kJ mol⁻¹ at 298 K (Figure S20), which rivaled the Gibbs free energy for the elongation of SP_{4Zn} (-38 kJ mol⁻¹, see above). In contrast, SP_{4Cu} was inert against DMAP because of the extremely low affinity of copper(II)-porphyrin for DMAP (Figure 5b)⁵⁷.



Figure 5. Kinetically enhanced stability of block supramolecular polymers. (a–d) Absorption spectra of SP_{4Zn} (a), SP_{4Cu} (b), a mixture of SP_{4Zn} and SP_{4Cu} (c), BSP($4c_u$ -*b*- $4z_n$ -*b*- $4c_u$) (d) with and without DMAP. (e) Time courses of absorbance changes because of the depolymerization of

SP_{4Zn} upon addition of DMAP: (i) SP_{4Zn}, (ii) SP_{4Cu}, (iii) a mixture of SP_{4Zn} and SP_{4Cu}, and (iv) BSP(4cu-*b*-4zn-*b*-4cu). [4zn] = [4cu] = 10 μ M and [DMAP, when added] = 200 μ M, in MCH/toluene (2:1) mixed solvent.

Interestingly, the depolymerization of SP4z_n by DMAP was prevented in the thus-synthesized BSP(4_{Cu}-*b*-4_{Zn}-*b*-4_{Cu}) structure ([4_{Zn}] = [4_{Cu}] = 10 μ M, Figure 5d,e and Figure S21). A small shoulder peak (430 nm) attributable to DMAP-4_{Zn} complex appeared mainly due to the coexisting monomeric 4_{Zn} in equilibrium (Figure 5d and see Figure S22 for detail). This result implied that the SP4_{Cu} blocks at both termini shielded the SP4_{Zu} block and prevented it from DMAP-triggered depolymerization. Importantly, the length histogram of the BSP(4_{Cu}-*b*-4_{Zn}-*b*-4_{Cu}) was retained even after the stability test (Figure S23).

For reference, the addition of DMAP to the mixture of SP_{4zn} and SP_{4cu} resulted in the selective depolymerization of SP_{4zn} (Figure 5c). Heating the solution of BSP(4_{Cu} -b- 4_{Zn} -b- 4_{Cu}) above T_e in the presence of DMAP and subsequent cooling led to a solution with an absorption spectrum identical to that obtained for the reference mixture of SP_{4zn}, SP_{4cu}, and DMAP (compare red lines in Figure 5c and d). In addition, application of sonication which breaks the BSP(4_{Cu} -b- 4_{Zn} -b- 4_{Cu}) thereby exposing the 4_{Zn} core to the DMAP also induced the same absorption spectral change (Figure S₂4). These results indicate that SP_{4zn} was kinetically stabilized against the DMAP ligand in the block structure.

We could also extend random SPs composed of 4zn and 4_{Cu} from the seeds of SP_{4Zn}. Depolymerization of the segment of SP_{4z_n} as an inner block was found to be stabilized against DMAP, suggesting that the blocks of random SP structure could also protect the inner \mbox{SP}_{4zn} (Figure S25). Previously, Schenning, Meijer, and coworkers^{54,56} reported the synthesis of a random SP of zinc(II)and copper(II)-porphyrin consisting monomers. They found that the zinc(II)-porphyrin monomer could be selectively removed from such a random SP by axial ligation with a Lewis base. A comparison of the kinetics of the system developed in the present study with Meijer's system may provide mechanistic insights into the depolymerization and monomer-exchange mechanisms.

Following previously reported methods for similar systems⁵⁸, molecular simulations was used in this study to obtain a molecular-level insight into these SPs and their DMAP-triggered depolymerization. All atom (AA) and coarse-grained (CG) models were developed for the 4_{Zn} monomer (Figure 6a). The CG scheme employed in this work was consistent with that recently used for 1,3,5-benzenetricarboxamide (BTA)-based SPs, allowing us to study the SPs at a resolution of ~5 Å⁵⁹. The AA model was used to fine-tune the CG model (see Method in Supporting Information). A pre-stacked SP model of the initially extended CG 4_{Zn} monomers was equilibrated in explicit cyclohexane solvent (equivalent to MCH at the level of resolution of these CG models) via CG molecular dynamics (CG-MD) simulation^{59,60} (here, we note that



Figure 6. Molecular modeling of 4_{Zn} supramolecular polymer. (a) Atomistic (AA) and coarse-grained (CG) models for the 4_{Zn} monomer. (b) Equilibrated configuration of the supramolecular fiber $4_{Zn}(CG)$ in cyclohexane: zinc(II)-porphyrin cores are colored in pink and the side chains are represented as grey/green transparent wires (solvent not shown for clarity). (c) AA and CG models for DMAP. Equilibrated configuration of fiber $4_{Zn}(CG)$ in cyclohexane (not shown) and DMAP (yellow). Only the DMAP molecules stably absorbed/interacting with the fiber are shown. Detail: DMAP coordination to zinc (in green); stable interactions are represented as green dynamic bonds. (d) Free energy (ΔG) for monomer exchange from the fiber tips (red arrow) obtained from WT-MetaD simulations. Data are shown as a function of the number of contacts between the exchanging monomer and the rest of the fiber (state A: assembled fiber; state B: exchange) for the $4_{Zn}(CG)$ fiber in cyclohexane in the absence (black) and presence of DMAP (red).

DMAP depolymerizes SP_{4zn} even in pure MCH). An equilibrated configuration of SP_{4zn} was obtained, which appeared as a very stable, extended fiber of unimolecular width under these conditions (Figure 6b). Next, explicit CG DMAP molecules were inserted in the simulation box with a DMAP:_{4zn} ratio of 20:1, and the system was re-equilibrated. It was observed that the DMAP molecules were adsorbed onto the surface of SP_{4zn} without being able to access the zinc atoms in the interior of SP_{4zn}. As shown in Figure 6c, it was only at the fiber tips that the DMAP molecules were stably coordinated to the porphyrin zinc atoms.

Next, the effect of DMAP on the depolymerization of SP_{4zn} was investigated. Following the recently reported method to study the dynamics of BTA-based SPs,⁶¹ welltempered metadynamics (WT-MetaD)⁶² simulation was used to activate and explore the monomer exchange process (see Method in Supporting Information). In the native condition (without DMAP), the global free energy difference between the assembled (A: N+1 assembled monomers) and disassembled state (B: N assembled + 1 disassembled) was found to be ~ 65 kJ mol⁻¹ (Figure S26). This value is in reasonable agreement with the experimentally determined ΔG for the elongation of SP_{4Zn} in MCH (52 kJ mol⁻¹, Figure S8). In the monomer exchange process from the SP_{4zn} tips, the system had to overcome a free energy barrier of \sim 75 kJ mol⁻¹ (Figure 6d, black). We also investigated monomer exchange process from the middle of SP_{4zn} via WT-MetaD simulations. In this mechanism, the SP_{4zn} first broke down to form a

defect or a new tip (Step 1: $\Delta G_1 = -70$ kJ mol⁻¹ needed), from which the monomer could then exchange (Step 2: see above): a mechanism also observed in BTA-based SPs in an organic solvent⁶¹. Assuming (underestimating) steps 1 and 2 as simply additive, and given the exponential relationship between the free energy barriers to cross and the characteristic probability for the events to occur, we can conclude that the monomer exchanges occurs predominantly from the fiber tips.

Interestingly, the same analysis on SP_{4zn} in the presence of DMAP demonstrated that the free energy barrier for monomer exchange from the fiber tip decreased to ~ 40 kJ mol⁻¹ (Figure 6d, red). Thus, free energy of the process was reduced by ~35 kJ mol⁻¹ when compared to the same SP in the absence of DMAP, which was in agreement with the competing DMAP-zinc experimental affinity in MCH (~30 kJ mol⁻¹: see, Figure S20). The free energy profile of monomer dissociation from the fiber tip obtained from WT-MetaD under such out-of-equilibrium conditions demonstrated that the DMAP molecules interfered with the monomer-monomer interactions at the fiber tip, first by penetrating in between the porphyrin cores and then by facilitating monomer detachment from the fiber tip (Figure S27). These results corroborate the stabilization scheme for BSPs demonstrated above, where SP_{4Cu} blocks at both tips of the SP_{4zn} block could prevent SP_{4zn} from DMAPtriggered depolymerization, by saturating the *de facto* depolymerization "hot spots" (i.e., the SP4zn tips). A similar strategy to enhance the stability of SP-one based

on the mechanism in which monomer exchange hot spots are decreased—has recently been reported by Palmans, Meijer, and co-workers⁶³ for supramolecular copolymers, but to the best of our knowledge, this is the first example demonstrated by using BSPs.

CONCLUSIONS

The concept of living supramolecular polymerization has rapidly evolved in recent years^{3,4,9-25}; nevertheless, the synthesis of BSPs still remains a challenge. As discussed at the beginning of this article, the dilemma of molecular design for connecting two distinct SPs needs to be addressed. Herein, it was demonstrated that BSPs could be synthesized in a similar manner as the living CDSA method^{16,17,34-37}. Although the conditions of seeded growth remain to be explored from the viewpoint of kinetics, the controlled length of the thus-obtained BSPs with narrow polydispersity confirmed that supramolecular polymerization occurred in the seeded growth mechanism.

It is noteworthy that SP_{4zn}, as an inner block in the BSP $(4_{Cu}-b-4_{Zn}-b-4_{Cu})$, could coexist with DMAP that was otherwise incompatible with SP4zn alone. Thus, it appears that a SP can be kinetically compartmentalized in the block structure. As illustrated in many biomolecular systems, compartmentalization is a key concept for achieving intricate multicomponent systems. The present finding can lead to many opportunities for the design of unprecedented functional supramolecular systems; in this context, for example, DMAP is a well-known organic catalyst and may function as such in the BSP matrix. Obviously, an emergence of unique properties from the combination and sequence of the blocks can also be expected⁶⁴. We believe that the synthetic strategy and characterization of BSPs demonstrated in this work will contribute to a better understanding of their attributes and applications for the development of multicomponent supramolecular systems.

ASSOCIATED CONTENT

Supporting Information. Experimental details, AFM images, spectroscopic data. "This material is available free of charge via the Internet at http://pubs.acs.org."

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Notes

The authors declare no competing financial interest.

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SYNOPSIS TOC

