1 Introduction

Supramolecular polymers, polymeric arrays of monomeric units that are brought together by reversible and highly directional secondary interactions, have been extensively applied for their versatility in the construction of intelligent artificial materials during the past few decades. Compared with traditional polymers, supramolecular polymers are endowed with many novel specific functions such as controllability, reversibility, responsiveness to external stimuli and ability to self-healing due to the dynamic and reversible nature of noncovalent interactions. With the further research, various noncovalent interactions have been widely attempted to facilitate the self-assembly of supramolecular monomers. For example, hydrogen bonding and metal–ligand interactions are usually employed as driving force for their high binding constants, macrocycle recognition and interlocked structures are frequently used as bridge to construct supramolecular polymers, etc. Supramolecular polymers can be functionalized with specific properties such as morphology adjustment, controllable luminescence, shape memory, self-healing, and so forth. Especially, the supramolecular polymers constructed in aqueous solution can offer more possibility for mimicking biocompatible or vital functional materials.

Cucurbit[8]uril (CB[8]) can accommodate two guest molecules in its cavity as a significant macrocycle. Scherman et al. reported the CB[8]-mediated heteroternary complexation with 1,1'-dimethyl-4,4'-bipyridinium dication (MV) and an uncharged azobenzene derivative to demonstrate the formation of CB[8] supramolecular polymers from guest monomers bearing azobenzene moieties both in solution and in the solid state. Zhang and co-workers reported a series of CB[8]-mediated supramolecular polymers with naphthalene- and anthracene-containing monomers. While there were rare reports about CB[8]-mediated heteroternary complexation with MV and coumarin derivative. And plentiful researches have indicated that short spacers are of high orientation selectivity which inhibits the formation of cyclic. Based on above researches, we designed and synthesized two multifunctional monomers which could assemble to form multi-morphology supramolecular polymers from linear to dendritic polymerization by the introduction of azobenzene in aqueous solution based on host–guest interaction. The first monomer is 1,1''-((hexane-1,6-diy1)bis(1-((2-oxy-2H-chromen-7-yl)methyl)-[4,4'-bipyridine]-1,1'-diium))bromide (HCDV; Scheme 1). HCDV contains coumarin–viologen–viologen–coumarin structure array with hexyl spacer as the flexible linkage which is suitable for supramolecular polymerization. The other is (E)-1,1''-(((diazone-1,2-diy1)bis(1-phenylene))bis(oxy))bis(butane-4,1-diy1)) bis(1-((2-oxy-2H-chromen-7-yl)methyl)-[4,4'-bipyridine]-1,1'-diium))bromide (PCDV; Scheme 1). PCDV is designed to comprising an azobenzene moiety in the middle of HCDV and contains two butyl spacers on either side of azobenzene as the flexible linkage. Moreover, we have

A novel supramolecular self-assembly based on ternary host–guest interaction between cucurbit[8]uril (CB[8]), 1,1'-dimethyl-4,4'-bipyridinium dication (MV) and coumarin derivative was applied for the construction of linear supramolecular polymer with high degree of polymerization in aqueous solution. Accompanied by the introduction of azobenzene on linear ABBA type monomer the supramolecular polymerization is different and the morphology changes from linear to dendritic polymer. The successful supramolecular polymerization of linear and dendritic supramolecular polymers by non-covalent host–guest molecular recognition was confirmed by various characterization methods, such as 1H NMR spectroscopy, ROESY, transmission electron microscopy (TEM) and dynamic light scattering (DLS) measurements. Meanwhile, the supramolecular polymerization could promote the conversion of the azobenzene from cis to trans, which ultimately results in no isomerism upon UV irradiation.
demonstrated that the coumarin was able to act as a second guest for a C8-MV binary complex, which led to the formation of a 1 : 1 : 1 heteroternary complex. Other reasons for choosing coumarin as a second guest include its high binding constant as well as its better optical and biological activity. The short length of flexible linkage of HCDV inhibits the formation of the 1 : 1 cyclic complex. Instead, the formation of linear supramolecular polymer is favored. The introduction of azobenzene in linkage site affects the polymerization pattern and thus forms dendritic supramolecular polymers. Apart from the transformation of polymerization pattern, the gradually introduction of C8 promotes the conversion of the azobenzene on the supramolecular polymer from cis to trans, which ultimately results in no isomerism upon UV irradiation. As far as we know, there were rare reports that azobenzene could be limited photoisomerization by supramolecular assemble methods.

2 Results and discussion

To confirm the formation of the ternary complex, 1H NMR experiments were performed in D2O (Fig. 1). After the addition of C8 to a solution of MV and 1-(2-oxo-2H-chromen-7-yl) methyl]pyridin-1-ium bromide (ChPy) with a molar ratio of 1 : 1 : 1, the 1H NMR signals of MV and ChPy shifted to upfield apparently with the signals for the protons in ChPy broadened. The process of self-assembly was also investigated by 1H NMR titration experiments which were performed in D2O by gradually increasing the ratio of ChPy to C8-MV (ESI, Fig. S1†). These changes indicated the successful encapsulation of MV and ChPy in the cavity of C8 and thus the formation of the ternary complex. A guest molecule 1-methyl-1-[(2-oxo-2H-chromen-7-yl)methyl]-[4,4'-bipyrindine]-1,1'-dium bromide (ChVio) in the cavity of C8 can be regarded as an AB-type monomer, with A and B representing viologen- and coumarin-moiety bound in one C8 respectively. Comparing the 1H NMR spectra for ChVio in the absence and presence of C8, the signals for protons in coumarin moiety still shifted to upfield and broadened. The peaks g, j, h, and i in viologen moiety also broadened besides shifting upfield which was presumably corresponding to polymeric species. These changes revealed that a linear supramolecular polymer with a low degree of polymerization may exist for a 1 : 1 mixture of C8 and ChVio.

The affinity between coumarin and C8-MV was obtained by isothermal titration calorimetry (ITC) experiments. Titration of ChPy into an aqueous solution of C8-MV gave an exothermic isotherm, and which indicated that C8-MV could bind with ChPy in a 1 : 1 molar ratio with binding constant K2 value of $6.67 \pm 1.22 \times 10^5$ M$^{-1}$ as shown in Fig. 1f. Such a high binding constant prefers to forming of a supramolecular polymer with a high degree of polymerization.

The linear supramolecular polymers with higher solubility were formed by mixing C8 and HCDV with a molar ratio of 2 : 1 based on the above assembly method. Furthermore, the introduction of azobenzene in the flexible linkage moiety generated another polymerization process and yielded dendritic supramolecular polymers whose schematic diagrams were shown in the Fig. 2. The stoichiometry between ChVio and C8, HCDV and C8 was confirmed to be 1 : 1 and 1 : 2 by their Job plots respectively, and the stoichiometry between C8 and PCDV was about 2.33 : 1 as shown in Fig. S6 in ESL†.
To confirm the formation of supramolecular polymers and study the assembly process of the supramolecular polymers, $^1$H NMR titration experiments were performed in D$_2$O. With the molar ratio of CB[8] to HCDV increased gradually, peaks of viologen and coumarin moieties in HCDV monomers broadened (Fig. S2 in ESI†), while the new peaks of the complex enhanced and the peaks of the free HCDV monomer disappeared when the molar ratio of CB[8]/HCDV was increased to 2 : 1 gradually. These changes suggested the formation of the supramolecular polymer in aqueous solution. Correspondingly, as the addition of CB[8] to PCDV, the signal H$_k$ in azobenzene moiety shifted upfield obviously, indicating that the azobenzene moiety was bounded in CB[8]. On the other hand, there was an obvious decrease in peak of free PCDV monomer with $\delta = 8.92$ when the molar ratio of CB[8]/PCDV was 1 : 1 (Fig. S3 in ESI†). When the molar ratio of CB[8]/PCDV exceeded 1.0, the peaks of the free PCDV monomer disappeared, which indicated that all PCDV monomers had been assembled to form the supramolecular polymers of CB[8] and PCDV. And when the molar ratio reached 2.0, all characteristic peaks broadened more and there were less sharp and dispersed signals in $^1$H NMR spectrum of PCDV-2CB[8] compared with the $^1$H NMR spectrum of HCDV-2CB[8]. Increased the molar ratio of CB[8] and PCDV to 2.5, the peaks of viologen and coumarin were further broadened, especially the coumarin moiety, which suggested that further supermolecular polymerization occurred. These analysis showed that there existed three assembly methods on PCDV/CB[8]. With the addition of CB[8] to PCDV, viologen and azobenzene moieties, viologen and coumarin moieties were bounded in the cavity of CB[8]. And when the molar ratio of CB[8] exceeded 2.0, two coumarin moieties were bounded in the cavity of CB[8]. The speculated process of the polymerization is depicted by a schematic diagram in the Fig. 2. On the other hand, 2D ROESY $^1$H NMR was employed to certify the interaction of proton signals in viologen and coumarin moieties, coumarin and coumarin$^\dagger$ as well as viologen and azobenzene (Fig. S4 and S5 in ESI†).

To further confirm the difference of supramolecular polymerization, transmission electron microscopy (TEM) measurements were undertaken. The TEM image of supramolecular polymer HCDV-2CB[8] and PCDV-2CB[8] gave distinctly different supramolecular self-assembled morphologies. As shown in Fig. 3a, an ultra-long curved string with more than a dozen micrometer length was observed, clearly indicating the existence of linear supramolecular polymer (for more detail morphology, see Fig. S8 in ESI†). By contrast, the TEM of PCDV-2CB[8] displayed dendritic morphology like “antler” shape with lengths of hundreds of nanometers to several micrometers in water (Fig. 3b). Such difference was consistent with the speculated polymerization process.

Dynamic light scattering (DLS) measurements in water were performed to provide the evidence for the existence of supramolecular polymer. A hydrodynamic radius of ca. 1.9 μm was found in HCDV-2CB[8], which indicated the formation of the supramolecular polymers with high degree of polymerization in water (Fig. 3c). By contrast, the PCDV-2CB[8] showed around 180 nm hydrodynamic size (Fig. 3d), but which was more evenly distributed. The results were consistent with the TEM and showed two different supramolecular polymerization process before and after the introduction of azobenzene. Meanwhile, the polymerization process of PCDV and CB[8] was also investigated by DLS. The results of PCDV-CB[8] and PCDV-2CB[8] showed that the hydrodynamic radius increased from 120 nm to 180 nm (Fig. S9†), indicating the increased degree of polymerization with the ratio of PCDV/CB[8] changed from 1.0 to 0.5. And the results were corroborated by our previous analysis of $^1$H NMR.

Meanwhile, the gradually introduction of CB[8] promotes the conversion of the azobenzene on the supramolecular polymer from cis to trans configuration, which ultimately results in no isomerism upon UV irradiation. The conversion process was investigated by UV/Vis spectroscopy. With the addition of CB[8] to PCDV, absorption peak at wavelength 366 nm decreased, accompanied by absorption peak at wavelength 450 nm increasing (Fig. 4a). A monomer PCDV could convert to cis from trans upon 365 nm UV irradiation. A reverse conversion happened as CB[8] was gradually added as the ratio of CB[8] to PCDV was increased from 0 : 3 to 6 : 3 (Fig. 4b). When the molar ratio of CB[8]/PCDV was 2 : 1, the supramolecular polymer PCDV-2CB[8] had no any response upon 365 and 254 nm (Fig. S10h in ESI†) UV irradiation. This phenomenon may be caused by strong affinity between CB[8] and two guest moieties limited isomerism of azobenzene in the middle, which was
Fig. 3  TEM images of the supramolecular polymer (a) HCDV-2CB[8] negative stained by Phosphotungstic acid and (b) PCDV-2CB[8] (40 uM) in water; DLS results of (c) HCDV-2CB[8] and (d) PCDV-2CB[8] (0.5 mM) in water.

Fig. 4  UV/Vis spectra of (a) PCDV (0.02 mM) and (b) under 365 nm UV irradiation with different ratios of CB[8] (PCDV/CB[8] = 3 : 0, 3 : 1, 3 : 2, 3 : 3, 3 : 4, 3 : 5, 3 : 6). Inset: magnification of the region 330–500 nm.
demonstrated by changing the molar ratio of CB[8] and PCDV irradiated on 365 nm UV for studying the conversion process (by UV/Vis spectroscopy shown in Fig. S10 in ESIF). And when the molar ratio of CB[8]/PCDV was smaller than 2:1, the photoisomerization happened to part of complex with the addition of CB[8] upon 365 nm UV irradiation, clearly indicating the photoisomerization of azobenzene limited by assembly of CB[8] and PCDV.

3 Conclusions

In summary, we demonstrated a novel supramolecular self-assembly based on ternary host-guest interaction between CB [8], MV and coumarin derivative and constructed a water soluble linear supramolecular polymer with high degree of polymerization. The higher binding constant between coumarin and CB[8]-MV was in favor of supramolecular self-assembly. Accompanied by the introduction of azobenzene on linear ABBA type monomer the supramolecular polymerization is different and the morphology changes from linear to dendritic polymer. Meanwhile, the supramolecular polymerization could promote the conversion of the azobenzene from cis to trans, which ultimately results in no isomerism upon UV irradiation. This research enriches the covalent interactions in the bottom-up nanofabrication of polymers and provides a significant route for controlling molecular self-assembly with tailored properties by non-covalent interactions in the bottom-up nanofabrication of molecular devices.

Conflicts of interest

There are no conflicts to declare.

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Notes and references


