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Structural effects of ditopic azoprobecyclodextrin complexes on the selectivity of guest-induced supramolecular chirality[†]

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Benzo-15-crown-5 and dipicolylamine are contained as the binding sites in a ditopic azoprobe (15C5-Azo-*n*-dpa). However, the selectivities of guest-induced supramolecular chirality for cations and anions were dramatically altered by a slight change in the spacer length of $(15C5-Azo-n-dpa)_2-\gamma-CyD$ complexes in water.

Chirality control by supramolecular assemblies and helical polymers based on chiral templates has attracted much attention in recent years.¹ Especially guest-induced chirality control is expected to apply for the development of versatile chiral switching and sensing systems.² To obtain the supramolecular chirality function, cyclodextrins (CyDs) are quite attractive host molecules.³ Optically inert CyDs have a chiral nature in their cavities and can be efficiently combined with various achiral chromoionophores and fluoroionophores to induce chiral nature by forming an inclusion complex with CyDs.⁴ In the previous study, we have reported a ditopic azoprobe (15C5-Azo-2-dpa) bearing benzo-15-crown-5 (B15C5) and dipicolylamine (dpa) as recognition sites. 15C5-Azo-2-dpa was found to form a 2:1 complex with γ -CyD and show a unique response function based on guest-induced supramolecular chirality in water.⁵ By allowing ditopic azoprobes to be incorporated into γ -CyD, we revealed the response behavior of the supramolecular $(15C5-Azo-2-dpa)_2-\gamma$ -CyD complex in the presence of each cationic and anionic species by measuring induced circular dichroism (ICD) spectra and UV-visible (Vis) absorption spectra. We confirmed that only when K^+ , Zn^{2+} , and CO_3^{2-} were all present, a large split-type Cotton effect appeared in the measured ICD spectra and a significant short-wavelength shift took place in the measured UV-Vis spectra. The result clearly demonstrates that the (15C5-Azo-2-dpa)₂- γ -CyD complex can exhibit supramolecular chirality due to the twisted

structure of the azoprobe dimer inside the γ -CyD cavity, only when it recognizes K⁺ and Zn²⁺ in the presence of CO₃^{2-,5} From the ICD spectral change, we can estimate the spatial changes of the azoprobe dimer inside γ -CyD, which induce a change in the UV-Vis spectra.⁶ Therefore each guest ion can be selectively detected in the presence of the other guest ions by measuring the spectral changes.

Herein we report how the **15C5-Azo-***n***-dpa** structure affects the selectivity of guest-induced supramolecular chirality. The dramatic selectivity changes of supramolecular chirality were found to be noted for (**15C5-Azo-***n***-dpa**)₂– γ -CyD complexes by controlling the spacer length of **15C5-Azo-***n***-dpa** from ethylene (*n* = 2) to propylene (*n* = 3) to butylene (*n* = 4) units (Fig. 1). Also, while **15C5-Azo-***n***-dpa** has a dpa binding site for heavy metal ions, the selectivity of guest-induced supramolecular chirality changed from Zn²⁺ for the (**15C5-Azo-2-dpa**)₂– γ -CyD complex to



Fig. 1 Structure of 15C5-Azo-n-dpa and the twisted structures.

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Cu²⁺ for the (**15C5-Azo-4-dpa**)₂– γ -CyD complex. However, most dpa-based chemosensors display Zn²⁺ and/or Cd²⁺ selectivity in water.⁷ Thus this is a unique example where the dpa-based supramolecular sensor exhibits a selectivity change from Zn²⁺ to Cu²⁺ caused by a change in the spacer length. We also report the specific selectivity changes of guest-induced supramolecular chirality for alkali-metal cations and anions based on the change in the spacer length of (**15C5-Azo-***n***-dpa**)₂– γ -CyD complexes in water.

The synthesis of **15C5-Azo-***n***-dpa** was carried out by the azocoupling of 4'-aminobenzo-15-crown-5 with phenol, followed by the introduction of bromoethylene, bromopropylene, and bromobutylene spacers using the Williamson ether synthesis.⁸ Then a dpa moiety was introduced under basic conditions with K_2CO_3 , and the obtained products were purified using silica gel column chromatography. The structures of **15C5-Azo-***n***-dpa** were confirmed *via* ¹H NMR and combustion analyses. Details of the synthesis are available in the ESI.[†]

Job's plot analyses clearly revealed that 15C5-Azo-n-dpa formed a 2:1 inclusion complex with γ -CyD in 4% DMSO-96% water (v/v) (Fig. S6, ESI⁺). The ICD spectra of (15C5-Azo-n-dpa)₂γ-CyD complexes are depicted in Fig. 2. For 15C5-Azo-2-dpa (Fig. 2a), the selective ICD response was only noted for Zn^{2+} over other metal ions (Mg2+, Fe3+, Ni2+, Cu2+, Cd2+, and Pb2+, as nitrate salts) in the presence of 50 mM K₂CO₃. The split ICD from the negative peak at 351 nm to the positive peak at 394 nm indicates the clockwise twisted structure of the two azoprobes inside the γ-CyD cavity.⁶ For 15C5-Azo-3-dpa, however, the ICD response was found to be observed not only for Zn²⁺ but also for Cu²⁺ in the presence of 50 mM K₂CO₃. Interestingly the ICD peak shape for Cu²⁺ was opposite compared with that for Zn²⁺, indicating that the Cu²⁺ complex formed an anti-clockwise twisted structure inside the γ -CyD cavity. This change in the twisted structure may be due to the difference in the coordination configuration; Cu²⁺ was capable of forming a coordination bond with the phenoxy ether oxygen in the dpa complexes,⁹ whereas only a few coordination bonds with the phenoxy ether oxygen were noted for the Zn²⁺-dpa complexes.¹⁰ For 15C5-Azo-4-dpa, the selective ICD response was only noted for Cu2+ over other metal ions (Mg²⁺, Fe³⁺, Ni²⁺, Zn²⁺, Cd²⁺, and Pb²⁺, as nitrate salts) in the presence of 50 mM K₂CO₃. Although 15C5-Azo-n-dpa possesses the same dpa binding site for heavy metal ions, the selectivity was dramatically changed from Zn²⁺ for the (15C5-Azo-2-dpa)₂-γ-CyD complex to Cu²⁺ for the (15C5-Azo-4-dpa)₂-γ-CyD complex in the presence of K₂CO₃. The (15C5-Azo-3-dpa)₂-γ-CyD complex exhibited selectivity for both Zn²⁺ and Cu²⁺, indicating an intermediate selectivity between 15C5-Azo-2-dpa and 15C5-Azo-4-dpa.¹¹ It is evident that the spacer length of 15C5-Azo-n-dpa played an important role in controlling the selectivity of guest-induced supramolecular chirality. As shown in Fig. 2, it should be noted that the shapes of split Cotton effects based on π - π * transition are not symmetric, indicating the overlap of the Cotton effect based on $n-\pi^*$ transition at the longer wavelength. In addition, the location of azobenzenes along the z-axis of CyD is known to strongly affect the sign and intensity of the Cotton effect.⁶ Although we consider that the bulky and hydrophobic B15C5 moieties restrict the movement of azobenzenes along the



Fig. 2 ICD spectra of **15C5-Azo-***n*-**dpa**/γ-CyD sensors in 4% DMSO aq.: (a) [**15C5-Azo-2-dpa**]; (b) [**15C5-Azo-3-dpa**]; (c) [**15C5-Azo-4-dpa**] = 0.04 mM, [Zn(NO₃)₂] = 0.04 mM, [Cu(NO₃)₂] = 0.04 mM, [γ-CyD] = 5 mM, [K₂CO₃] = 50 mM.

z-axis of the γ -CyD cavity, the abovementioned factors make the detailed understanding of guest-induced ICD responses difficult. To obtain further evidence for the guest-induced ICD responses, additional analysis based on molecular mechanics and TD-DFT calculations as well as X-ray crystallography analysis are to be conducted.

In the presence of equivalent amounts of Zn^{2+} with **15C5**-**Azo-***n***-dpa** (20 μ M), and 50 mM CO₃²⁻, the ICD intensities at the maximum wavelength are plotted against the alkali metal ion diameter (Fig. 3a). As we reported previously, the (**15C5-Azo-2dpa**)₂- γ -CyD complex exhibited high K⁺ ion selectivity over other alkali metal ions in the presence of Zn^{2+} and CO_3^{2-} . This selectivity is consistent with the selectivity of sandwich complex formation of the two benzo-15-crown-5 derivatives with alkali metal ions.^{8,12} However, for the (**15C5-Azo-3-dpa**)₂- γ -CyD complex, the alkali metal ion selectivity was significantly reduced (Fig. 3a). This indicates that the formation of the clockwise twisted structure is dominated only by the bridge formation of CO_3^{2-} with the two dpa-Zn²⁺ complexes in the



Fig. 3 Selectivity of **15C5-Azo-n-dpa**/ γ -CyD sensors toward alkali metal ions in 4% DMSO aq., [γ -CyD] = 5 mM, [alkali metal ion] = 50 mM: (a) [**15C5-Azo-2-dpa**], [**15C5-Azo-3-dpa**] = 0.04 mM, [Zn(NO₃)₂] = 0.04 mM; (b) [**15C5-Azo-3dpa**], [**15C5-Azo-4-dpa**] = 0.04 mM, [Cu(NO₃)₂] = 0.04 mM.

(15C5-Azo-3-dpa)₂– γ -CyD complex. The enhanced flexibility of 15C5-Azo-*n*-dpa upon changing the spacer from ethylene (n = 2) to propylene (n = 3) should be the reason of this selectivity change. On the other hand, in the presence of equivalent amounts of Cu²⁺ with 15C5-Azo-*n*-dpa (20 µM), and 50 mM CO₃²⁻, the (15C5-Azo-3-dpa)₂– γ -CyD complex exhibited high K⁺ ion selectivity similar to the Zn²⁺ system of the (15C5-Azo-2-dpa)₂– γ -CyD complex (Fig. 3b). However, for the (15C5-Azo-4-dpa)₂– γ -CyD complex, no alkali metal ion selectivity was observed in the anti-clockwise twisted structure of the two azoprobes inside the γ -CyD cavity (Fig. 3b). This is also ascribed to the enhanced flexibility of 15C5-Azo-*n*-dpa upon changing the spacer from propylene (n = 3) to butylene (n = 4).

The effect of anion species on the ICD responses was also examined for $(15C5-Azo-n-dpa)_2-\gamma$ -CyD complexes. For the Zn²⁺ complex system of $(15C5-Azo-n-dpa)_2-\gamma$ -CyD complexes (n = 2, 3), the effects of salt species on ICD responses were examined in the presence of 50 mM KX (X = NO₃⁻, CH₃CO₂⁻, and OH⁻). The results are depicted in Fig. 4. Similar to the $(15C5-Azo-2-dpa)_2-\gamma$ -CyD complex, the $(15C5-Azo-3-dpa)_2-\gamma$ -CyD complex exhibited high CO₃²⁻ selectivity, indicating that CO₃²⁻ bridging with the two Zn²⁺-dpa binding sites induced the clockwise twisted structure of the azoprobe dimer inside the γ -CyD cavity. In addition, direct evidence of the relative orientation of the azoprobes and the macrocyclic ring was obtained *via* NOESY experiments. Cross-peaks between H3 protons inside the CyD cavity and protons of the azoprobes were clearly observed (Fig. S7, ESI⁺).



Fig. 4 Selectivity of (**15C5-Azo-n-dpa-Zn²⁺**)₂- γ -CyD sensors toward anions in 4% DMSO aq.: (a) [**15C5-Azo-2-dpa**] = 0.04 mM; (b) [**15C5-Azo-3-dpa**] = 0.04 mM, [Zn(NO₃)₂] = 0.04 mM, [γ -CyD] = 5 mM, [Cl⁻], [OH⁻], [NO₃⁻], [CH₃CO₂⁻], [CO₃²⁻] = 50 mM.







On the other hand, for the Cu²⁺ complex system of (15C5-Azo-*n*-dpa)₂- γ -CyD complexes (n = 3 and 4), the (15C5-Azo-*n*-dpa)₂- γ -CyD complexes showed both CO₃²⁻ and OH⁻ selectivity (Fig. 5). This indicates that hydroxo-bridging between the two Cu²⁺-dpa binding sites induced the anti-clockwise twisted structure of the azoprobe dimer in the (15C5-Azo-*n*-dpa)₂- γ -CyD complex. Thus, by changing the metal species of the dpa binding sites, the anion selectivity of the (15C5-Azo-3-dpa)₂- γ -CyD complex can be easily controlled from the CO₃²⁻ selectivity with the Zn²⁺ system to both CO₃²⁻ and OH⁻ selectivity with the Cu²⁺ system in water. These are apparently a unique function of guest-induced supramolecular chirality based on (15C5-Azo-*n*-dpa)₂- γ -CyD complexes.

In conclusion, we have shown a novel guest-induced supramolecular chirality induced by twisted structural switching of the two **15C5-Azo-***n***-dpa** molecules inside the γ -CyD chiral cavity due to multi-point recognition of guest ions by the ditopic azoprobes in water. Although the two recognition sites are the same, a slight change in the spacer length of **15C5-Azo-***n***-dpa** was found to significantly affect the ICD response selectivity of (**15C5-Azo-***n***-dpa**)₂- γ -CyD complexes. To the best of our knowledge, this is a novel selectivity control based on guestinduced supramolecular chirality which completely differs from the design strategy of conventional molecular recognition systems.

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Conflicts of interest

There are no conflicts to declare.

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