

Stimuli-induced folding cascade of a linear oligomeric guest chain programmed through cucurbit[*n*]uril self-sorting ($n = 6, 7, 8$)[†]

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A six-station linear guest for cucurbit[7]uril and cucurbit[8]uril has been synthesized in order to implement a cascade of transformations driven by external stimuli. The guest chain is sequence-programmed with electron-deficient viologen and electron-rich naphthalene stations linked by either flexible or rigid spacers that affect the chain's folding properties. Together with the orthogonal guest selectivity of the two cucurbiturils, these properties result in self-sorted cucurbituril pseudorotaxane foldamers. Each transformation is controlled by suitable chemical and redox inputs and leads not only to refolding of the guest chain, but also to the liberation of secondary messenger molecules which render the system presented here reminiscent of natural signaling cascades. The steps of the cascade are analyzed by UV/Vis, ¹H NMR and electrospray (tandem) mass spectrometry to investigate the different pseudorotaxane structures in detail. With one guest oligomer, three different cucurbiturils, and several different chemical and redox inputs, a chemical system is created which exhibits complex behavior beyond the chemist's paradigm of the pure chemical compound.

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Introduction

Systems chemistry is a new emerging field in chemistry¹ which breaks with the current paradigm of the pure, isolated compound which has been one of the foundations of synthetic chemistry for decades. In contrast, systems chemistry aims at complex mixtures of compounds which cooperate synergistically in reactivity networks. The metabolism of a living cell offers the role model for such a chemical network. It does not only produce and metabolize the compounds needed for homeostasis and reproduction, but also regulates itself in spatiotemporal patterns of utmost complexity far away from thermodynamic equilibrium. Regulation often involves signal transduction through membranes so that outer stimuli can affect the living cell to react to its environment. But also inside a cell, signaling cascades operate in which enzymes refold in conformation upon a messenger binding to a certain receptor site and thus inducing an active conformation that generates secondary messengers.

A concise description of such a network would require the detailed knowledge of the differential equations describing the concentration changes for all metabolites involved. Even then, the pattern evolution can only be modeled numerically.

Feedback loops remain problematic; in particular as self-accelerating processes² tend to amplify small initial deviations from a given set of starting parameters put into such a calculation. As a major consequence, predicting the long-term behavior of a complex chemical system is at least difficult, if not impossible. The more so, this is true for the *de novo* design of synthetic chemical networks with a certain desired behavior.

Another interesting conceptual idea evolves, if one accepts that there is no fundamental difference between living and inanimate matter: evolution is then not restricted to the biological world anymore, but can become an emergent property³ of a synthetic system as well.⁴ Consequently, the motivation to investigate chemical networks comes not only from the desire to emulate and understand nature,⁵ but also from the interest in creating chemical systems to put our conceptions on evolution and emergence to the test – or to create systems that have so far new and unseen properties.

Several different approaches have led to systems displaying emergent properties, for example self-sorting in supramolecular systems,⁶ the adaptive behavior of dynamic combinatorial libraries (DCLs),⁷ self-replicators that feed on DCLs,⁸ or mechanosensitive supramolecular fiber formation.⁹ Recently, Nitschke *et al.* reported metallo-supramolecular systems which are responsive to external stimuli and undergo cascading transformations that are reminiscent of signaling cascades in living cells.¹⁰

During the last decades, the host-guest chemistry of the cucurbit[*n*]uril family¹¹ ($n = 5-8, 10$) has seen tremendous interest mainly because of strong binding constants,¹² a wide

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variety of guests, the cavity-size-dependent differences in guest preferences,¹³ and the potential to implement switching behavior.¹⁴ Only recently, few groups have been focusing on the dynamic self-sorting of these hosts.¹⁵ Previously, we reported a detailed study on the social self-sorting of cucurbit[7]uril (CB7) and cucurbit[8]uril (CB8) in the presence of guests like viologen, dihydroxy naphthalene and the corresponding covalently linked di- and trimers of both.¹⁶ As a result, specific supramolecular architectures form in this social network, because CB7 binds viologen dications, while CB8 prefers guest pairs of electron-deficient viologen and electron-rich dihydroxy naphthalene.¹⁷ One of the still quite rare cases has been discovered, in which *both* hosts CB7 and CB8 have a preference for the *same* guest A over B, but still socially self-sort when stoichiometry is controlled, because the binding constants of the CB7·A and CB8·A pairs are sufficiently different. Besides pairs of electron-deficient and electron-rich guests, CB8 also strongly binds dimers of viologen cation-radicals.¹⁸ This motif has found applications in the design of redox-stimulus-driven molecular

machines,¹⁸ but no studies are available that discuss how the cation-radical dimer affects the self-sorting behavior of CB7 and CB8. Consequently, cucurbiturils have large potential for studying their behavior in complex chemical systems.

Here, we present the five-step transformation cascade shown in Fig. 1. The cascade involves a linear guest chain **BATCl₈** with six stations which allow inducing the formation of different foldamers by different signals added to the guest. Elements of the underlying program include the different cavity sizes and thus different guest selectivities of the three different cucurbit[n]urils CB6–CB8 ($n = 6–8$), the sequence of stations along the guest chain and the rigidity/flexibility of the spacers connecting them. This transformation cascade is thus reminiscent of nature's signaling cascades in that external signals cause recognition events that lead to controlled conformational changes in a sequence-programmed chain and result in the liberation of secondary messenger molecules.

Results and discussion

Concept and guest chain synthesis

To implement the stimuli-induced folding processes depicted in Fig. 1, the guest molecule **BATCl₈** is designed to have a specific sequence of guest units: two terminal viologens (T) are connected to the adjacent naphthalenes through flexible propyleneoxy linkers; the two inner viologens (I) are linked to the naphthalenes through more rigid triazolyethylene spacers. The central connection between the two inner viologens is again a flexible propylene group. The propyleneoxy linkers are known to be the shortest linkers permitting the viologen–naphthalene guest dimer to fold into a charge-transfer (CT) complex of the electron-poor viologen and the electron-rich naphthalene units.¹⁹ The central propylene linker instead helps maximizing the formation of the cofacial viologen cation-radical dimer upon two one-electron reductions at the central viologen moieties.²⁰ In contrast, suitable folding into CT complexes of the central viologen and the naphthalene units is made impossible by the more rigid triazolyethylene spacers. Thus, the guest chain involves two design elements: the sequence of binding stations and the incorporation of suitable spacers, which either allow or prohibit folding of the chain, respectively.

BATCl₈ is synthesized in a convergent manner by initially connecting the two terminal viologen–naphthalene pairs (see ESI† for details). For solubility, the azide-substituted hydroxy-naphthalene is first equipped with a bromopropylene side chain and then reacted with *N*-methyl-4,4'-bipyridinium iodide. For the synthesis of the inner part of the guest, 4-bromo-1-butyne is reacted with 4,4'-bipyridine. Before connecting the two inner viologens to each other, an anion exchange of the bromide to the corresponding hexafluorophosphate is necessary for solubility reasons. After this exchange, the two inner viologens are connected with 1,3-dibromopropane and the product subsequently equipped with the two terminal parts through a copper-catalyzed Huisgen–Sharpless–Meldal click reaction. In order to obtain the final, water-soluble chloride salt, another anion exchange is performed.

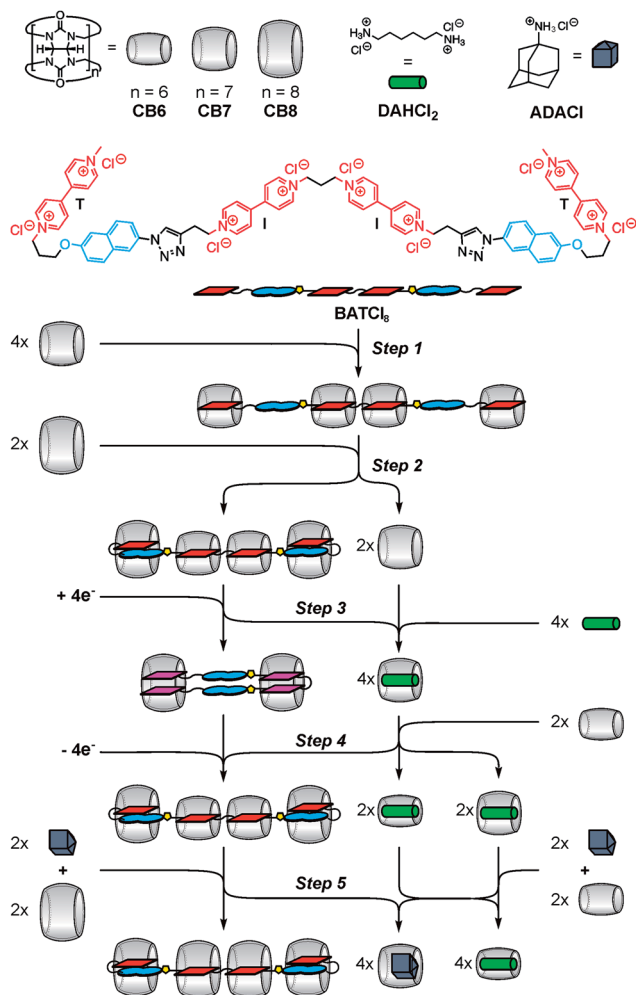


Fig. 1 Top: the cucurbiturils and guest cations used in this study and the cartoons used to represent them. Bottom: the five-step transformation cascade in which different external signals lead to adaptive responses of the host–guest complexes.

Characterization of the cascade intermediates

With each step of the transformation cascade, new external stimuli are added either as chemical or redox signals. Consequently, the mixture becomes more and more complex in each step. Classical NMR analysis of complexation-induced signal shifts can provide very useful structural information on the host-guest complexes formed. However, the interpretation of the NMR spectra might become increasingly difficult with the complexity of the mixture under study. A second complementary method is thus advantageous to cross check the results obtained from NMR experiments.

The BAT^{8+} guest chain is – as well as the other guests DAH^{2+} and ADA^+ – charged and thus facilitates a mass-spectrometric analysis of the mixtures obtained after each step in the cascade. This method has been utilized before and showed that even different isomeric host-guest complexes can be distinguished by their fragmentation patterns.¹⁶ Therefore, (tandem) electrospray Fourier-transform ion-cyclotron-resonance mass spectrometry (ESI-FTICR-MS) is used in the present study as one of the major methods for the characterization of the different host-guest complexes formed in each of the transformation steps along the cascade. There are three peculiarities in the gas-phase behaviour of viologens and their cucurbituril complexes, which are important for understanding the corresponding mass spectra: (i) the naked viologen dications are prone to fragmentation induced by charge repulsion unless stabilized by host-guest interactions or the presence of counterions.²¹ (ii) When cucurbiturils are bound to the viologens, most often the highest possible charge states are observed with remarkably high intensity. This is in contrast to the free guests, which usually appear in lower charge states because they carry a number of counterions.¹⁶ This finding can be rationalized by the increase in anion-cation distance during complexation of the cucurbituril. It is thus by far easier to strip off the counterions from the cucurbituril complexes than from the free guests upon ionization. Nevertheless, charge-state distributions are often observed. (iii) Viologen dications and cation-radicals sometimes undergo one-electron reduction or oxidation reactions at the ESI capillary so that charge states can be observed other than those expected from the number of counterions still present in the complexes.

In addition, UV/Vis spectroscopy provides insight into the formation of the viologen cation-radicals and their dimerization products upon reduction of BATCl_8 . These experiments are thus helpful, when NMR experiments are hampered by the paramagnetism of the viologen cation-radicals.

First step: complex formation of BATCl_8 with CB7

The first step of the transformation cascade is the formation of a 4 : 1 complex of CB7 and BATCl_8 . The binding constant of methyl viologen to CB7 has been reported²² to amount to $K = 2.0 \times 10^5 \text{ M}^{-1}$, a value certainly high enough to justify the expectation that this complex will form. It is simply generated by adding 4 eq. of CB7 to a 0.857 mM water solution of BATCl_8 . After equilibration, the expected host-guest complex $[\text{BAT@CB7}_4]^{8+}$ is detected as the base peak in the electrospray

Fourier-transform ion-cyclotron-resonance (ESI-FTICR) mass spectra at m/z 738 (Fig. 2, center). In addition, the $[\text{BATCl@CB7}_4]^{7+}$ (m/z 849) and $[\text{BATCl}_2\text{@CB7}_4]^{6+}$ (m/z 996) ions are observed with lower intensities. Also traces of $[\text{BAT@CB7}_5]^{8+}$ (m/z 884) are detected, which is likely an unspecific complex as often detected in ESI mass spectra. This assignment is supported by a fragmentation experiment (see ESI†). Only those fragments are observed that are also found for $[\text{BAT@CB7}_4]^{8+}$. Consequently, the fifth CB7 is only very weakly bound and easily stripped off the complex. Finally, the $[\text{BAT@CB7}_3]^{8+}$ ion at m/z 593 is generated by loss of a CB7 from $[\text{BAT@CB7}_4]^{8+}$. One viologen is not stabilized by the cucurbituril anymore and thus a charge separating fragmentation into the fragments $[\text{A@CB7}]^{3+}$ (m/z 528) and $[\text{B@CB7}_2]^{5+}$ (m/z 626) occurs easily. Consequently, this mass spectrum is in good agreement with expectation for the formation of $[\text{BAT@CB7}_4]^{8+}$.

When mass-selected $[\text{BAT@CB7}_4]^{8+}$ ions are fragmented with a CO_2 IR laser in infrared multiphoton dissociation (IRMPD) experiments (Fig. 2, bottom), three competing pathways are observed in a rather complex fragmentation behavior. The loss of one CB7 with subsequent dissociation into $[\text{A@CB7}]^{3+}$ and $[\text{B@CB7}_2]^{5+}$ is the first and likely least energy-demanding one of them. The second fragmentation channel is

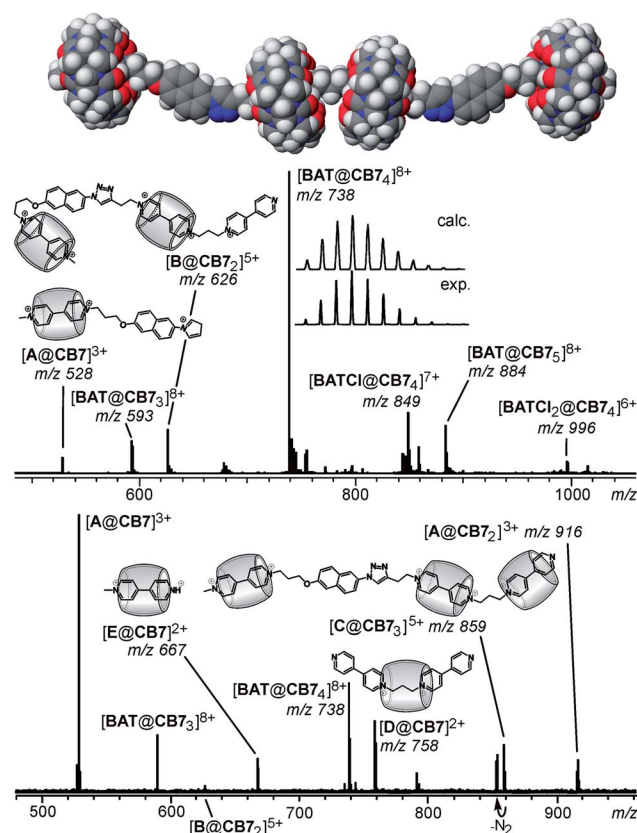


Fig. 2 Top: force-field optimized structure (MM2, CaChe 6.1 program, Fujitsu, Krakow, Poland) of the $[\text{BAT@CB7}_4]^{8+}$ complex. Center: electrospray FTICR mass spectrum of the 0.857 mM sample solution of BATCl_8 and 4 eq. of CB7 in water (after dilution to ca. 30 μM). Bottom: infrared multiphoton dissociation experiment with mass-selected $[\text{BAT@CB7}_4]^{8+}$ ions. Also, see ESI† for details.



a charge-separating dissociation into fragments $[A@CB7]^{3+}$ and $[C@CB7_3]^{5+}$ (m/z 859). The third reaction is cleavage of the parent ion into $[B@CB7_2]^{5+}$ and $[A@CB7_2]^{3+}$ (m/z 916). The $[C@CB7_3]^{5+}$ ion undergoes CB7 and nitrogen molecule²³ losses followed by a fragmentation into $[A@CB7]^{3+}$ and $[D@CB7]^{2+}$ (m/z 758). Finally, $[E@CB7]^{2+}$ (m/z 667) is the likely end point of several fragmentation pathways.

This analysis of the reaction pathways is based on the experience that a multiply charged ion preferentially fragments into two smaller daughter ions, which are both charged and thus should both appear in the spectrum. With this prerequisite, a number of other potential channels can safely be ruled out as only one of the expected product ions is observed. The fact that cucurbituril loss competes with covalent bond cleavages is indicative of the formation of rather strong supramolecular host-guest interactions and thus points to the formation of a specific 4 : 1 complex.

The 1H and $^1H,^1H$ COSY NMR spectra (see ESI†) of this complex reveal the two outer cucurbiturils to reside almost centrally on the terminal viologens. However, the CB7 molecules bound to the inner viologens appear with two sets of signals. The less intense one corresponds to CB7 located centrally on the inner viologens. The more intense one can be assigned to a CB7 that is shifted towards the triazole ring. As this structure is the major species in the equilibrium, it is likely that the triazole C–H hydrogen forms a C–H \cdots O hydrogen bond²⁴ with one of the carbonyl groups in the CB7 outer rim as also indicated by the significant downfield shift of the triazole C–H signal ($\Delta\delta \approx 0.5$ ppm). Similar “half-threaded” complexes have been observed earlier.²⁵ Consequently, the NMR spectra are in line with the formation of a 4 : 1 complex and add more structural information in that they show isomeric complexes to exist in an equilibrium that is on slow exchange with respect to the NMR time scale.

Second step: competition between CB7 and CB8

In the second step of the transformation cascade, 2 eq. of CB8 are added in competition to the already present CB7 hosts. CB8 binds methyl viologen with a binding constant similar to that of CB7 ($K = 1.1 \times 10^5 M^{-1}$).²⁶ With its larger cavity, it can also very favorably bind a charge-transfer guest pair. For example, the binding constant of the methyl viologen–dihydroxynaphthalene pair amounts to $K = K_1 \times K_2 \approx 10^9 M^{-2}$.²⁷ Earlier studies¹⁶ revealed a clear self-sorting phenomenon: when an electron-deficient viologen and an electron-rich naphthalene are connected covalently, CB8 binds significantly stronger to this pair than a CB7 to the viologen alone. Based on this self-sorting behavior, the addition of CB8 to the sample solution obtained in the first step should lead to the formation of the $BATCl_8@CB7_2 \cdot CB8_2$ complex with a well-defined CB8-CB7-CB7-CB8 sequence. The external CB8 signal is thus expected to give rise to a refolding of the guest chain together with the liberation of two molecules of CB7.

Clear-cut evidence for this successful transformation comes from ESI mass spectrometry as well as NMR spectroscopic experiments (Fig. 3). The ESI-FTICR mass spectrum of the

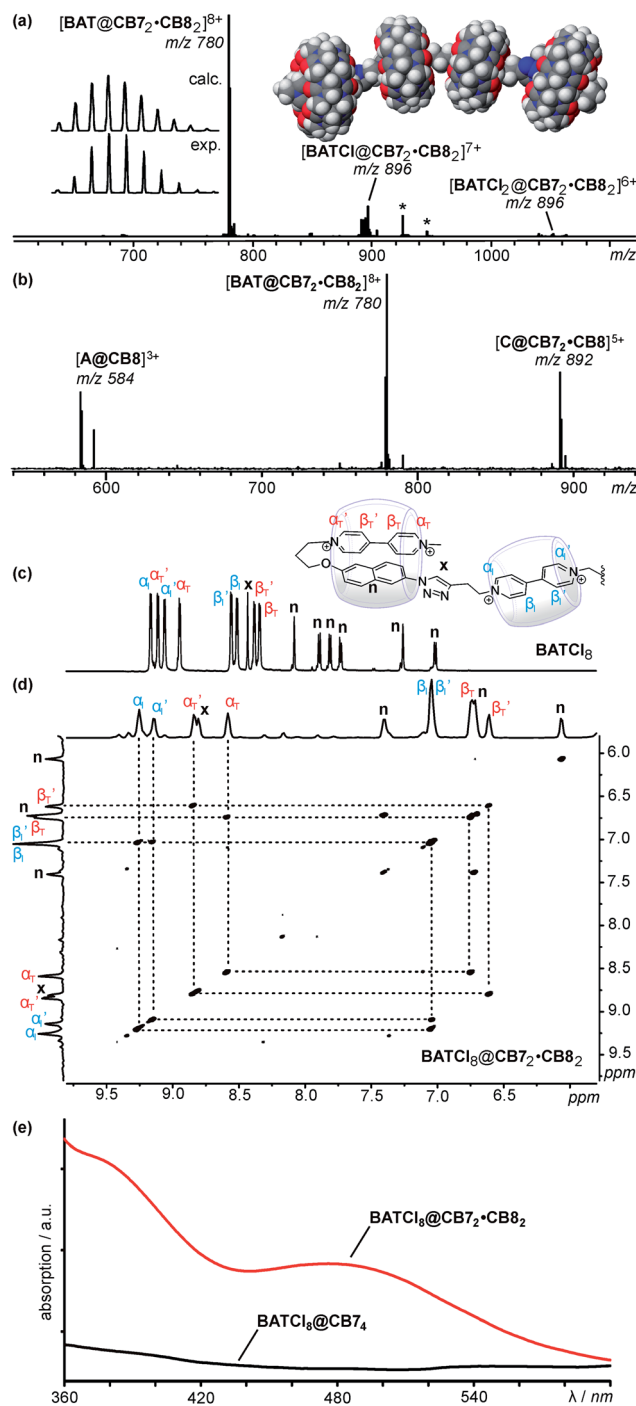


Fig. 3 (a) ESI mass spectrum of a sample solution obtained after addition of 2 eq. of CB8 to the 0.857 mM sample from step 1 followed by dilution to ca. 30 μM . Asterisks indicate signals due to unspecific complex formation with CB7 or CB8, respectively. Inset: force-field optimized structure of $[BAT@CB7_2 \cdot CB8_2]^{9+}$. (b) IRMPD experiment performed with mass-selected $[BAT@CB7_2 \cdot CB8_2]^{8+}$ ions. (c) 1H NMR spectrum (700 MHz, D_2O , 298 K) of a 0.857 mM water solution of the free guest chain $BATCl_8$. (d) $^1H,^1H$ COSY NMR spectrum of a 0.857 mM water solution of $BATCl_8$ in presence of 4 eq. of CB7 and 2 eq. of CB8. (e) UV/Vis absorption spectra of 1.22×10^{-2} mM water solutions of $BATCl_8@CB7_2 \cdot CB8_2$ (red) and $BATCl_8@CB7_4$ (black).



sample after **CB8** addition exhibits only one major signal which can be assigned to the 1 : 2 : 2 complex $[\text{BAT}@\text{CB7}_2 \cdot \text{CB8}_2]^{8+}$ (m/z 780). Again, smaller signals for the next two lower charge states are observed as well as two unspecific complexes, which carry either a **CB7** or a **CB8** in excess. So far, the spectrum is analogous to that of the $[\text{BAT}@\text{CB7}_4]^{8+}$ complex discussed above. However, one remarkable difference is obvious: there are no fragments formed that could be traced back to the loss of a cucurbituril. This can be attributed to the stronger binding of the two **CB8** molecules, which act as stoppers trapping the two inner **CB7** molecules. This finding already provides a first indication of the cucurbituril positions along the chain.

These considerations are supported by the IRMPD experiment (Fig. 3b) performed with mass-selected $[\text{BAT}@\text{CB7}_2 \cdot \text{CB8}_2]^{8+}$. In contrast to that done with $[\text{BAT}@\text{CB7}_4]^{8+}$, only a single fragmentation pathway is observed – a charge separating cleavage into $[\text{A}@\text{CB8}]^{3+}$ (m/z 584) and $[\text{C}@\text{CB7}_2 \cdot \text{CB8}]^{2+}$ (m/z 892). No loss of **CB8** competes with this covalent bond cleavage reaction. This points again to the significantly enhanced binding strength of **CB8**. More importantly, no $[\text{A}@\text{CB7}]^{3+}$ fragment ion is observed thus providing evidence for the relative positioning of **CB7** in the center and **CB8** at the termini of the guest chain.

A comparison of the ^1H NMR spectra of the free host BATCl_8 and the $\text{BATCl}_8@\text{CB7}_2 \cdot \text{CB8}_2$ complex (Fig. 3c and d) reveals the expected complexation-induced shifts. All viologen β protons reside inside hydrophobic cavities and experience upfield shifts of about 1.5 ppm.²² The same holds true for the naphthalene protons so that we can safely conclude the naphthalene units to be located inside the cavities of the two **CB8** molecules. The viologen α protons do not move as much as they are located at the rims of the cucurbiturils. While the α_1 protons for the inner viologens experience a minor downfield shift, the α_T protons of the terminal viologens are shifted somewhat upfield indicating that they are affected by the anisotropy of the adjacent naphthalene. Consequently, the NMR shifts are structure-indicative and confirm the terminally folded 1 : 2 : 2 complexes of BATCl_8 and the two cucurbiturils. Again, the triazole C–H signal is shifted downfield indicating that C–H \cdots O hydrogen bonds are also formed in this complex. In contrast to $\text{BATCl}_8@\text{CB7}_4$, however, no positionally shifted isomers are observed anymore. We conclude the **CB8** host molecules to bind to the triazole C–H with one of their carbonyl oxygen atoms. Therefore, the **CB7** molecules now prefer to be located centrally on the inner viologens and do not shift towards the triazole anymore.

Finally, the formation of the viologen–naphthalene charge-transfer complexes inside the **CB8** cavities is also confirmed by the presence of the two typical absorption bands at 383 nm and 488 nm in the UV/Vis spectrum of the complex (Fig. 3e).¹⁹ These two bands are absent in $\text{BATCl}_8@\text{CB7}_4$ and only appear in the presence of **CB8**.

All these results provide a picture not only consistent with the formation of a folded $\text{BATCl}_8@\text{CB7}_2 \cdot \text{CB8}_2$ complex with **CB8** in the peripheral and the **CB7** in the central positions along the guest chain. It also nicely agrees with the details observed in the first step of the transformation cascade. The differences between both complexes in the tandem MS fragmentation and NMR experiments provide quite detailed insight into the

structures of both complexes. It should be noted that the second step does not only comprise reorganization of the host–guest complex and refolding of the chain into a more contracted structure, but also the liberation of two empty **CB7** molecules reminiscent of the formation of secondary messengers in signaling cascades in nature.

Third step: refolding through viologen cation-radical dimerization inside **CB8**

Initially, the third step was intended to lead to yet another foldamer of the guest chain using a redox instead of a chemical signal. It is well known that viologen dications can be reduced to the corresponding cation-radicals, which tend to dimerize in particular, when **CB8** is present providing stabilization to the dimer by encapsulation. However, the first attempts to achieve a four-electron reduction of the guest chain with Zn powder failed and only incomplete reduction was obtained (see ESI†). Therefore, $\text{Na}_2\text{S}_2\text{O}_4$ has been used as the reductant. But still, the presence of free **CB7** in the sample solution caused incomplete refolding of the chain as **CB7** binds to the viologen cation-radical monomer and thus competes with **CB8** efficiently (see ESI†).²⁸ Finally, the addition of the reductant together with 4 eq. 1,6-diaminohexane dihydrochloride DAHCl_2 as a scavenger for all **CB7** resulted in a clean transformation. The third step thus requires the simultaneous action of a redox and a chemical signal.

When the sample solution obtained in step 2 is treated with 20 eq. of $\text{Na}_2\text{S}_2\text{O}_4$ as the reductant and 4 eq. of DAHCl_2 , the ESI mass spectrum in Fig. 4a is obtained after a reaction time of 24 hours. Clearly, signals appear for the $[\text{DAH}@\text{CB7}]^{2+}$ dication (m/z 641) accompanied by two signals in which one or two protons have been exchanged against sodium ions (m/z 652 and 663). The second species detected is indeed the quadruply reduced $[\text{BAT}@\text{CB8}_2]^{4+}$ complex (m/z 978). As discussed above, one-electron oxidation reactions can occur at the ESI needle in the ion source. This would be one potential reason, why the $[\text{BAT}@\text{CB8}_2]^{5+}$ ion (m/z 782) is also observed. A second possibility is of course the incomplete reduction of the viologens.

To distinguish both possibilities, the reduction was investigated by UV/Vis spectroscopy. First, an independent sample containing BATCl_8 and 2 eq. of **CB8** was prepared in water. Directly after the addition of $\text{Na}_2\text{S}_2\text{O}_4$, the UV/Vis spectrum shows the presence of both the viologen cation-radical dimer with its typical bands at 367, 509, 538, and 865 nm and the viologen cation-radical monomer as indicated by the additional bands at 394 and 601 nm.¹⁴ Over time, the bands of the monomer decrease, while the ones of the dimer increase to reach complete conversion after ca. 2 hours. Usually, the dimerization of the cation-radical is a very fast process proceeding in a millisecond time regime.¹⁹ A control experiment with free BATCl_8 confirmed this to be true for this guest, too. Immediately after the addition of the reductant, only cation-radical dimers were observed in the UV/Vis spectra (see ESI†). We therefore interpret our results for the $\text{BATCl}_8@\text{CB8}_2$ complex as follows: before the reduction of the guest chain, a 2 : 1 complex of **CB8** and BATCl_8 forms. As only the terminal viologens can form CT



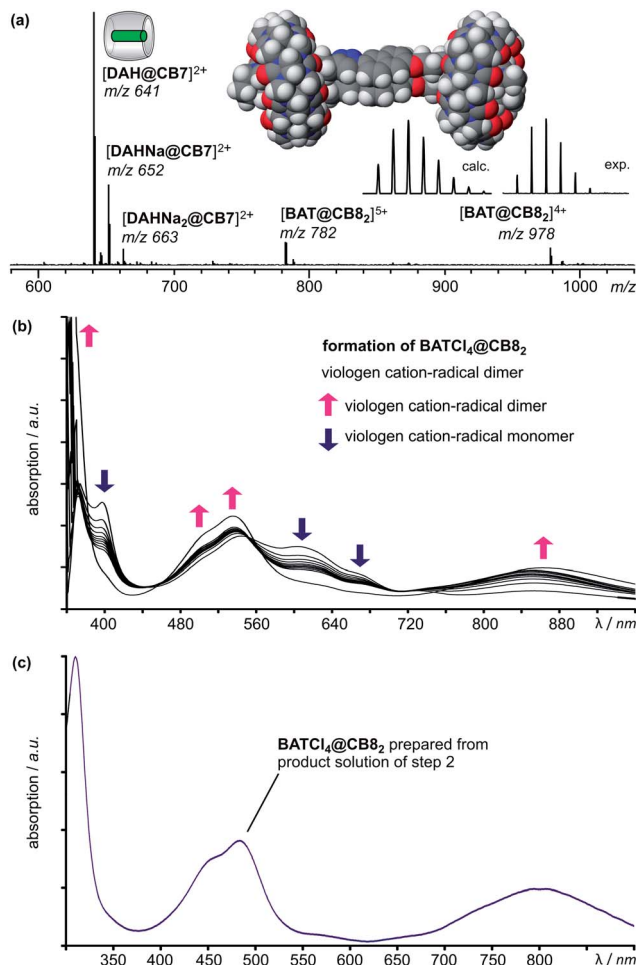


Fig. 4 (a) ESI mass spectrum of a sample solution obtained after reduction using 20 eq. of $\text{Na}_2\text{S}_2\text{O}_4$ and addition of 4 eq. of DAHCl_2 to the sample from step 2 (then diluted to ca. 30 μM). Inset: force-field optimized structure of the $[\text{BAT@CB8}_2]^{4+}$ complex. (b) Changes in the UV/Vis spectra upon $\text{Na}_2\text{S}_2\text{O}_4$ (20 eq.) reduction of an independently prepared 8.90×10^{-2} mM water solution of BATCl_8 containing 2 eq. CB8 (reaction time: two hours). (c) UV/Vis spectrum of the final product of the transformation of the sample from step 2 after the addition of 20 eq. $\text{Na}_2\text{S}_2\text{O}_4$ and 4 eq. DAHCl_2 and a reaction time of 24 hours.

complexes with the adjacent naphthalenes, the CB8 molecules will certainly be located at the termini. Directly after the addition of reductant, the two inner viologens are quickly reduced and dimerize giving rise to the dimer bands in the UV/Vis spectra already right at the beginning of the experiment. The terminal viologens are, however, protected against reduction by CT complex formation and its stabilization by CB8 binding. This reduction proceeds thus much more slowly and stepwise so that the cation-radical monomer is observed as an intermediate. In the end, the monomers recombine into the folded structure shown in Fig. 4a, which contains two cation-radical dimers, both of which bind one CB8 .

The UV/Vis spectrum of $\text{BATCl}_4@CB8_2$ prepared by fully reducing a sample solution obtained from the second step of the cascade (figure c) clearly reveals the exclusive presence of cation-

radical dimers, while the typical bands of the monomers are absent. Consequently, we conclude not only the $[\text{BAT@CB8}_2]^{5+}$ ion in the ESI mass spectra to be generated during the electrospray process, but the third transformation step to proceed cleanly. The cooperation of a redox and a chemical signal give rise to yet another foldamer of the guest chain.

Forth step: going a step back by reoxidation and the addition of cucurbit[6]uril

Reoxidation of the guest chain is possible by bubbling air through the sample solution over a period of 10 minutes and the UV/Vis spectrum indicates that the cation-radical dimers are back-converted into viologen dications. Also, the bands typical for the charge-transfer complexes of the terminal viologens and the naphthalenes reappear (see ESI†). The simultaneous addition of 2 eq. of cucurbit[6]uril scavenges half of the DAHCl_2 guests, which bind more strongly to CB6 than to CB7 . Consequently, 2 eq. of CB7 become available again to reform the $\text{BATCl}_8@CB7_2\cdot\text{CB8}_2$ complex.

The ESI mass spectrum in Fig. 5a clearly exhibits signals for all expected complexes and provides evidence for the simultaneous formation of $\text{DAHCl}_2@CB6$ and $\text{DAHCl}_2@CB7$ together with $\text{BATCl}_8@CB7_2\cdot\text{CB8}_2$, while other potentially possible complexes are not observed. The exclusive formation of these three complexes is also supported by the $^1\text{H}, ^1\text{H}$ COSY NMR spectra (see ESI†), which very clearly show the formation of the viologen–naphthalene charge transfer complexes incorporated in $\text{BATCl}_8@CB7_2\cdot\text{CB8}_2$ as well as the formation of both $\text{DAHCl}_2@CB6$ and $\text{DAHCl}_2@CB7$ in equal amounts. The

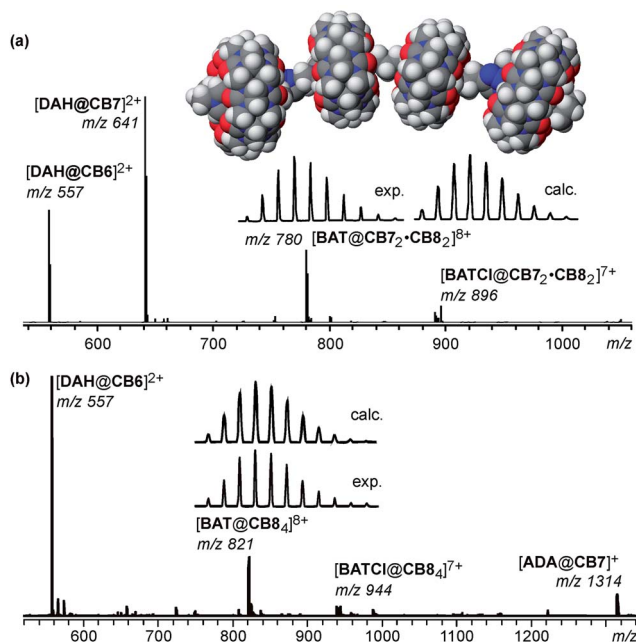


Fig. 5 (a) ESI mass spectrum of the sample solution after reoxidation of the guest chain and CB6 addition in the fourth step. (b) ESI mass spectrum after the addition of CB6 , CB8 , and ADACl in the fifth step. Again, the sample solutions used for electrospray ionization were diluted to ca. 30 μM before being admitted to the ion source.

chemical shifts observed for the latter two complexes are in agreement with those reported earlier.^{13a}

Fifth step: a complicated cucurbituril “metathesis”

Finally, the aim of the last step in the cascade reported here aims at the exchange of the two CB7 molecules located at the central viologens against two slightly weaker binding CB8 molecules to produce the $\text{BATCl}_8\text{@CB8}_4$ complex. This goal can be achieved, when 2 eq. of CB8 are added simultaneously with 4 eq. ADACl as a preferred guest for CB7. As this will liberate two DAHCl_2 guest dications from the remaining $\text{DAHCl}_2\text{@CB7}$ complexes, it is furthermore required to add 2 eq. of CB6 in order to obtain a clean transformation.

Fig. 5b shows the corresponding ESI mass spectrum, which – in view of the many different compounds now present in the mixture – is surprisingly clean and nicely confirms the self-sorting to work as expected as long as stoichiometry is carefully controlled. In the mass spectrum, DAHCl_2 is only observed in a complex with CB6 (as $[\text{DAH@CB6}]^{2+}$ at m/z 557) and ADACl only binds to CB7 (as $[\text{ADA@CB7}]^+$ at m/z 1314). Besides these two ions, the only more intense ion corresponds to $[\text{BAT@CB8}_4]^{8+}$. Again, ^1H , ^1H COSY NMR spectra support these mass spectrometric results (see ESI†). While the signals for $\text{DAHCl}_2\text{@CB6}$ prevail, those observed in step 4 for $\text{DAHCl}_2\text{@CB7}$ vanish and new signals for ADACl@CB7 appear.^{13a} Also, the typical chemical shifts of the viologens and naphthalenes forming the charge transfer complex are present. In conclusion, also the last transformation step in the cascade has successfully been carried out.

Conclusions

A transformation cascade has been presented which is reminiscent of natural signaling cascades although it only uses synthetic molecules. Nevertheless, conceptually, there are similarities: (i) a sequence-programmed guest chain is used instead of a biopolymer with sequence information. (ii) External signals result in molecular recognition events that at least in some of the steps of our cascade lead to conformational changes in the guest chain. Different foldamers are generated depending on the external stimuli. (iii) In some steps, secondary messengers are liberated that bind other molecules present in the mixture. As long as the stoichiometry is balanced, the self-sorting works nicely. (iv) Quite complex mixtures are generated, in which the guest selectivities of the three different cucurbiturils synergize with the sequence programmed into the guest chain and its rigidity-mediated folding properties to give rise to a surprisingly clear-cut behavior.

Certainly, there are also significant differences between natural signalling cascades and that presented here. (i) While in our case, the system always finds its way back into thermodynamic equilibrium after addition of each of the chemical stimuli, natural cascades are kept far away from thermodynamic control by an energy flow through the living organism. (ii) Natural systems contain feedback loops. Together with the energy flow, which drives them, emergent properties arise that

are much more complex than those observed here. (iii) Although our system is much less complex than natural systems, it is likely more sensitive to deviations from ideal stoichiometries.

Our study is a proof-of-principle study which paves the way into much more complex systems in the future. Many other steps can be imagined that could be performed with the system presented here. The diversity increases even more, when one considers guest chains with different and maybe longer sequences. This clearly indicates the potential of our approach for future work.

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