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A supramolecular keypad lock†

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The reversible photoswitching between an anthracene derivative and its [4+4] dimer, using the template effect of the CB8 macrocycle, was demonstrated. This example of supramolecular chemistry in water was harnessed to demonstrate the operation of a keypad lock device that is driven by means of light and chemicals as inputs.

The use of molecular species for achieving information processing of chemical, photonic or electrochemical signals according to the principles of binary coding (0 and 1) and Boolean language continues to receive wide attention.^{1–3} The challenge of achieving the design of molecules that are capable of a predetermined logic operation has received important backup from smart applications of molecular logic gates in sensing, drug delivery, theranostics, and materials chemistry.4-10 Among the different logic operations, those that imply a memory function belong to the most demanding ones in terms of their chemical design.¹¹ This type of logic, known as sequential logic, results in differentiated outputs depending on the input history of the device. Current research activities along these lines have their focus on the implementation of flip-flops and molecular keypad locks. Photochromic switches have been often exploited for these purposes.¹²⁻¹⁵ However, chemically-addressable systems and redox-switchable molecular devices were also used for the demonstration of molecular keypad locks¹⁶⁻¹⁹ and flip-flops.^{20,21}

Surprisingly, the exploitation of supramolecular host-guest phenomena for the demonstration of keypad lock functions has no precedence in the literature. In recent reports host-guest complexes with cucurbit[7]uril (CB7) and cucurbit[8]uril (CB8)

^a CIQSO - Center for Research in Sustainable Chemistry and Department of Chemical Engineering, Physical Chemistry and Organic Chemistry, University of Huelva, Campus El Carmen s/n, E-21071 Huelva, Spain. were used to demonstrate the reconfigurable and resettable operation of logic gates in aqueous solution.^{22,23} Cucurbiturils have drawn much attention for their very high binding constants of cationic guests (up to 10^{17} M^{-1})²⁴ and their supramolecular application potential with a strong focus on biological and pharmacological contexts, and analytical problems is beginning to reveal.^{25–35}

In the present work we take advantage of the specific complexation properties of the CB8 macrocycle. Unlike the smaller homologues CB6 and CB7, that commonly offer space for only one guest molecule, the larger CB8 often accommodates two guests and the resulting complexes may feature new emission properties (*e.g.*, excimer fluorescence) or lead to fluorescence self-quenching.^{35–38} Additionally, the resulting pre-organization of the two guests may facilitate intracomplex photoreactions that would not happen at the dilute concentrations of the free dye molecules.^{38–40} In a wider context, host-templated photodimerizations have been used in the design of photoswitchable supramolecular polymers.^{40–42}

Herein we designed the anthracene derivative **1** (see the ESI[†] for details of the synthesis and Scheme 1 for the structure) which contains a positively charged tetraalkylammonium side chain, known to interact efficiently with the carbonyl-lined portals of CB8. The aromatic parts of the guests should be immersed in the hydrophobic inner cavity of the host macrocycle leading to a 2:1 (guest:host) complex. This situation can give rise to a template effect in a [4+4] photodimerization,⁴⁰



Scheme 1 Reversible photoswitching between the anthracene derivative 1 and its photodimer 1₂; structures of CB7 and CB8.

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a well-known photoreaction of anthracenes.⁴³ The ammonium arms are expected to be oriented in opposite directions occupying the two portals of the macrocycle and thereby avoiding destabilization of the complex by electrostatic repulsion of the positive charges. This binding mode and the long-axis symmetry of **1** would lead to a single dimerization product (**1**₂; see Scheme **1**). Drawbacks for photoswitching are undesired secondary photoreactions that can hamper the efforts to achieve complete reversibility by cycloreversion of the photodimer.⁴⁰ Photooxidation reactions of anthracenes may interfere, especially when working with highly energetic UV light in aerated solutions. This obstacle can be potentially avoided by using electron-poor aromatic systems such as dye **1**.

With these design outsets we started the characterization of the supramolecular interactions between dye 1 and CB8. In the titration of a 10 µM aqueous solution of 1 with CB8 at neutral pH significant changes in the UV/vis absorption spectrum were observed (see Fig. 1a). The characteristic long-wavelength absorption of free 1 at λ_{max} = 412 nm is bathochromically shifted. The uniformity of the interaction was confirmed by the occurrence of various isosbestic points at 319, 349, and 424 nm. The absorption titration curve levelled-off sharply at ca. 0.5 equivalents of CB8, pointing to the predicted formation of the 1.1.CB8 complex (see ESI[†]). This stoichiometry was independently verified by Job's method (see ESI[†]). In electrospray-ionization mass spectrometry the existence of the 2:1 complex was also corroborated for the gas phase (see ESI[†]). Furthermore, ¹H NMR studies evidenced pronounced upfield shifts of the aromatic proton signals of the guest dye, coinciding with its deep immersion into the CB8 macrocycle (see ESI[†]). The fluorescence of **1** (λ_{max} = 494 nm, $\Phi_{\rm f}$ = 0.44, $\tau_{\rm f}$ = 6.72 ns) underwent significant variations in the course of the titration with the CB8 host, leading to the observation of a strong emission quenching; see Fig. 1b. This is explained by an enhanced non-radiative deactivation caused by π - π interactions between the face-to-face organized aromatic guest molecules. The fitting of the titration data gave a binding constant of K = $4.5 \times 10^{12} \text{ M}^{-2}$, which is indicative of a strong supramolecular



Fig. 1 Spectral changes of the UV/vis absorption (a) and the fluorescence (b) upon titration of 1 (3.6 μ M) with CB8 (0–6.2 μ M) in pH-neutral aqueous solution.



Fig. 2 UV/vis absorption spectral changes upon irradiation of **1** (10 μ M) in the presence of CB8 (5 μ M) at λ > 395 nm in pH-neutral aqueous solution. The inset shows the kinetic curves and the mono-exponential fittings for the irradiation of **1** in the absence (empty circles) and the presence of CB8 (filled circles); observation at 412 nm (free dye) or 424 nm (presence of CB8).

interaction and the avoidance of electrostatic repulsion of the positively charged arms of **1** (see above).

Having established the strong binding of 1 to CB8, the intracomplex photodimerization of the dye was attempted. Irradiation (150 W Xe-lamp, $\lambda_{exc} > 395$ nm) of a diluted solution of 1 (10 µM in neutral water) in the presence of CB8 (5 µM) resulted in a uniform photoreaction that was monitored by UV/vis absorption spectroscopy, liquid chromatography, and NMR spectroscopy (see ESI⁺). Under the applied conditions ca. 75% of the long-wavelength absorption band (between ca. 350 and 500 nm) of the complex depleted within 30 min of irradiation; Fig. 2. In comparison, the free dye (*i.e.*, in the absence of CB8) showed only ca. 10% decrease of the longwavelength absorption band under identical irradiation conditions (inset of Fig. 2). Hence, a much faster photoreaction (by a factor of ca. 10) resulted in the presence of CB8. By means of liquid chromatography with mass spectrometric detection the CB8-templated formation of the dimerization product was unequivocally confirmed by the observation of a molecular ion peak at m/z = 347 with an isotope pattern spacing of $\Delta m =$ 0.5 amu (see ESI⁺). ¹H NMR studies provided structural proof for the formation of photodimer 1_2 . Namely, pronounced changes in the aromatic region and a new signal at ca. 4.4 ppm, indicative of the bridgehead protons of the dimer, were detected (see ESI⁺). On the other hand, no photodimer was noted for the irradiation of the noncomplexed dye at $\lambda > 395$ nm (see ESI[†]).[‡] The cycloreversion of the photodimer $\mathbf{1}_2$ (see Scheme 1) back to the monomeric dye 1 was achieved by irradiation at 254 nm for a short period of time (30 s). The efficient back reaction was signalled by the recovery of more than 90% of 1 as rated using UV/vis absorption spectroscopy. Also in the ¹H NMR spectra the recovery of the aromatic signals that correspond to the complexed monomeric dye and the concomitant disappearance of the respective photodimer signals were evident (see ESI[†]). The recycling of the dimerization/cycloreversion sequence was shown for at least 5 cycles, submitting the CB8 complexes to successive irradiation at >395 nm and 254 nm (see ESI⁺).



Scheme 2 Supramolecular keypad lock by applying the correct input order, reading the output with the help of the "developer input", and resetting by CB7.

The described CB8-template effect on the photodimerization of 1 as well as the reversible photoswitching forms the chemical basis for the realization of the announced supramolecular keypad lock (see Scheme 2). For this purpose the inputs were chosen as the irradiation at $\lambda > 395$ nm (input A) and the addition of the strongly binding competitor 1-aminoadamantane (input B).^{35,44} It is noteworthy that the latter is able to displace the monomeric dye $\mathbf{1}$ as well as the dimer $\mathbf{1}_2$ from the CB8 macrocycle. The output of the system was defined as the observation of noncomplexed photodimer. The detection of this situation would in principle demand a cumbersome analysis, for example, using NMR spectroscopy. In order to visualize the output situation more conveniently, the irradiation at 254 nm ("developer input" in Scheme 2) together with the resulting change of the fluorescence intensity ($\Delta I = I_{254} - I_0$; the subscripts 254 and 0 refer to after and before irradiation at 254 nm, respectively) was monitored. The photodimer $\mathbf{1}_2$ is non-fluorescent both in the absence and presence of CB8. However, the cycloreversion of the noncomplexed dimer back to 1 is accompanied by a recovery of the fluorescence, while the same process inside CB8 leads to no significant fluorescence increase due to efficient quenching (see ESI[†]). In accordance, the only high output in form of ΔI (being defined as binary 1 and corresponding to the situation of the open lock) was observed for the sequential input order of first A and then B. This coincides with the formation of the noncomplexed photodimer (see Scheme 2). All other combinations led to no photodimer (input combinations 00, B0, 0B, BA) or to the formation of photodimer residing inside the CB8 cavity (A0 and 0A), see Fig. 3.

The resetting of the system to its initial state would be a valuable asset. The reversible nature of the photoreaction and of the involved supramolecular interactions enable such a feature. Taking for example the situation of the open keypad lock (illustrated in Scheme 2), the competitor 1-aminoadamantane is encapsulated by CB8 ($K = 3.3 \times 10^7 \text{ M}^{-1}$)³⁵ and dye **1** is a



Fig. 3 Operation of the supramolecular keypad lock (10 μ M **1**, 5 μ M CB8) upon application of the corresponding inputs (A: irradiation at λ > 395 nm for 45 min; B: addition of 1-aminoadamantane; 20 μ M). The output is read as the change of the fluorescence upon irradiation at 254 nm for 30 s. The dotted line shows the threshold. With respect to input A, stronger light sources would enable the shortening of the irradiation time. Note that the input combinations also include that one or both inputs may not be activated (binary 0).

noncomplexed species after application of the "developer input" (see Scheme 2). Given the fact that the binding constant of 1-aminoadamantane to CB7 is three orders of magnitude larger $(K = 1.2 \times 10^{10} \text{ M}^{-1})^{35}$ than for CB8 and that dye **1** has a much smaller binding constant with CB7 ($K = 3.0 \times 10^5 \text{ M}^{-1}$; this work), thermodynamic self-sorting can be used to achieve resetting.44 Indeed, the addition of CB7 to the mixture that was obtained after irradiation at 254 nm led to the complexation of 1-aminoadamantane by CB7 and of 1 by CB8, thereby closing the cycle shown in Scheme 2. The spectral signature of the 1.1.CB8 complex was re-constituted by addition of CB7 (see ESI[†]). At this point the logic operations can be repeated. This was shown for 5 cycles, after which a fatigue effect of ca. 30% was noted (see ESI⁺). A concentration optimization of the 4 components that are involved in the resulting intricate multi-equilibrium situation⁴⁵ should improve the performance further.

In conclusion, the anthracene dye **1** forms a reversibly photoswitchable complex with CB8, harnessing the template effect exerted by the organic macrocycle.⁴⁰ This was used to demonstrate a supramolecular keypad lock where the formation of a noncomplexed photodimer, indirectly signalled by the fluorescence enhancement upon photoinduced cycloreversion, was the output. Only by application of the right input order (first irradiation at $\lambda > 395$ nm and then addition of 1-aminoadamantane) the lock was opened. By exploitation of self-sorting upon addition of CB7 the system was reset to the initial 1·1·CB8 complex. These results demonstrate the utility of cucurbituril chemistry for the realization of resettable complex logic operations with all-organic systems in aqueous solution. Applications as functional switches in bio-relevant contexts are foreseen.

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Notes and references

‡ Applying the Stern–Volmer formalism and the fluorescence lifetime of **1** (see text) leads to the conclusion that even a diffusion-controlled [4+4] photodimerization (k_{diff} ca. 10¹⁰ M⁻¹ s⁻¹) of noncomplexed **1** would be very inefficient (<1%) in diluted solution (10 μ M).

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