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## **PAPER**



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# Remote electron and energy transfer sensitized photoisomerization of encapsulated stilbenes†

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Excited state chemistry and physics of molecules, in addition to their inherent electronic and steric features, depend on their immediate microenvironments. This study explores the influence of an organic capsule, slightly larger than the reactant molecule itself, on the excited state chemistry of the encapsulated molecule. Results presented here show that the confined molecule, in fact, is not isolated and can be manipulated from outside even without direct physical interaction. Examples where communication between a confined molecule and a free molecule present outside is brought about through electronic and energy transfer processes are presented. Geometric isomerization of octa acid encapsulated stilbenes induced by energy and electron transfer by cationic sensitizers that attach themselves to the anionic capsule is examined. The fact that isomerization occurs when the sensitizer present outside is excited illustrates that the reactant and sensitizer are communicating across the molecular wall of the capsule. Ability to remotely activate a confined molecule opens up new opportunities to bring about reactions of confined radical ions and triplet excited molecules generated via long distance energy and electron transfer processes.

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## Introduction

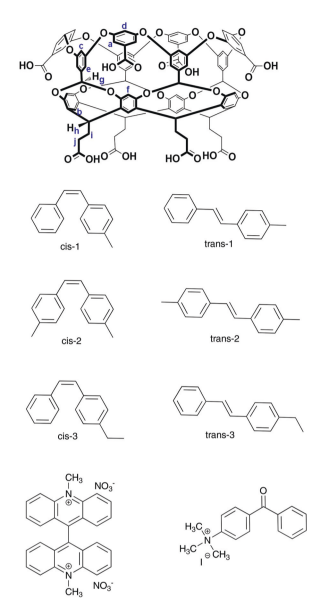
It is a well-known fact that phototransformations of organic molecules can be brought about by direct light absorption and/or through indirect excitation via sensitization. 1-3 In the second process, a second molecule known as a sensitizer absorbs the incident light and triggers energy and/or electron transfer to generate the triplet of the reactant or radical ion pairs, respectively. During the last six decades, the phenomena of triplet-triplet energy transfer and electron transfer processes have been extensively investigated and their value in chemistry, biology and materials science has been established.1-5 During the last decade, propelled by the demand to devise 'new green synthetic methods', photochemistry, especially energy and electron transfer processes under the name of 'visible light photocatalysis' (VLPC), has witnessed a surge in activity. 6-10 For over three decades, we have been interested in manipulating photochemical reactions employing supramolecular concepts. 11-19 Envisaging that blending the concepts of supramolecular chemistry (SC)<sup>19</sup> and VLPC<sup>20</sup> would yield new opportunities in constructing organic mole-

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cules in a selective fashion, we have explored the possibility of conducting energy and electron transfer between a confined reactant molecule and a free sensitizer that is present outside the confined space. Furthermore, ability to manipulate encapsulated molecules from outside opens up a new dimension in supramolecular photochemistry. In this context, we have recently established the occurrence of spin, energy and electron transfer between confined and free molecules through time resolved photophysical techniques.<sup>21–30</sup> These examples unequivocally established that communication between molecules occurs even if one of them is incarcerated in a molecular capsule. In this article, we have exploited the remote activation concept to bring about photoreactions of encapsulated molecules by activating them through energy and electron transfer sensitization from outside. In this study, the supramolecular host is a capsule made up of two molecules of a synthetic cavity and octa acid (OA),31 the guests are stilbenes, the electron transfer sensitizer is bis-N-methylacridiniumnitrate (BMAN, also known as lucigenin), the triplet energy transfer sensitizer is N,N,N-trimethyl-4-(phenylcarbonyl)benzenaminium iodide (4-TMABP) and the photoreaction used as the probe is the well-known geometric isomerization around C=C bonds (Scheme 1).32-34

Mechanistic details of electron and energy transfer sensitized geometric isomerization of stilbenes are well documented. 32,33,35-38 The triplet energies of cis and trans-stilbenes are closer to 57 and 49 kcal mol<sup>-1</sup>, respectively, 33 while oxidation potentials of these isomers are in the range of 1.75



**Scheme 1** Chemical structures of host octa acid (OA), guest stilbenes and sensitizers *-N*-methylacridiniumnitrate (BMAN) and *N,N,N*-trimethyl-4-(phenylcarbonyl) benzenaminium iodide (4-TMABP).

to 2.05 eV (Ag/AgI) with  $cis \sim 0.1$  eV higher than the corresponding trans. The energy transfer sensitizer 4-TMABP with triplet energy >60 kcal mol<sup>-1</sup> should be able to sensitize both isomers of stilbenes while the electron transfer sensitizer with a reduction potential of -0.3 eV (Ag/AgCl in water) and an excitation energy of  $\sim 2.7$  eV should be able to oxidize both isomers. Results that establish the occurrence of electron and triplet energy transfer sensitized geometric isomerization of OA encapsulated stilbenes 1, 2 and 3 with BMAN and 4-TMABP as sensitizers are discussed in this presentation. One should note that while the triplets of stilbenes isomerize both ways (cis to trans and trans to cis),  $^{35}$  the radical cations of stilbenes generated by electron transfer isomerize only one way,  $^{36}$  cis to trans. Therefore, the triplet sensitization experiments

with 4-TMABP were performed with both *cis* and *trans* isomers while electron transfer sensitization with BMAN was conducted only with the *cis* isomer.

## Experimental

#### **General information**

All commercially available chemicals were used as supplied without further purification, unless otherwise noted. <sup>1</sup>H NMR characterization, NMR titration studies and diffusion experiments were performed on a Bruker Avance 500 spectrometer equipped with a cryoprobe. Deuterated solvent was used as a lock and a residual protonated solvent peak was used as a reference. Absorption spectra were recorded on a Shimadzu UV-3150 spectrophotometer. Steady-state luminescence spectra were recorded using a FS920CDT fluorometer (Edinburgh Analytical Instruments).

#### Synthesis of host and guest molecules

Host octa acid (OA) was synthesized and characterized according to the reported procedure.<sup>31</sup> The stilbenes were synthesized according to the literature procedures.<sup>36,39</sup> Bis-*N*-methylacridiniumnitrate was purchased from Aldrich and used as received.

#### Synthesis of 4-TMABP

Methyl iodide (0.93 mL, 1.5 mmol) was added dropwise to a stirred solution of 4-aminobenzophenone (10 mmol) in DMF (5 mL) at room temperature. The mixture was stirred for 24 h at 50 °C. Following this, the reaction mixture was cooled to RT and diethyl ether (15 mL) was added dropwise to the solution. The resulting mixture was kept aside for 12 hours and filtered. The residue was washed with ether and dried, and crystallized at room temperature from methanol to afford 4-TMABP as a pale gray powder. A pure product (490 mg, 25%) was obtained and characterized by <sup>1</sup>H, <sup>13</sup>C NMR and mass spectrometry.

<sup>1</sup>H NMR (500 MHz,  $D_2O$ ): δ 7.97 (s, 4H), 7.79 (d, 2H), 7.72 (t, 1H), 7.56 (t, 2H), 3.67 (s, 9H).

<sup>13</sup>C NMR (125 MHz, D<sub>2</sub>O):  $\delta$  198.93, 149.54, 138.68, 135.97, 134.11, 132.02, 130.37, 128.73 120.20, 56.97.

HRMS (ESI): calculated for  $C_{16}H_{18}NOI^+$  [M - I] $^+$  240.14, found 240.1394.

#### Complexation of guest molecules

To a  $D_2O$  (10 mM borate buffer) solution of OA (1 mM) 5  $\mu L$  of 60 mM (DMSO-d<sub>6</sub>) of guest solution was added in gradual increments. After each addition, the NMR spectrum was recorded.

#### Identification of the location of the sensitizers

To locate the cationic sensitizers BMAN and 4-TMABP in the presence of stilbene@OA<sub>2</sub>, buffer solutions containing the capsules of cis-1@(OA)<sub>2</sub> and trans-1@(OA)<sub>2</sub> were prepared by adding 5  $\mu$ L of 60 mM respective standard guest solution to 0.6 mL of 1 mM of OA in 10 mM of borate buffer solution. To

this, 60 µL of 20 mM of BMAN or 4-TMABP solution were added in a step wise manner. After each addition, the NMR spectrum was recorded.

#### Irradiation

For electron transfer experiments, a solution containing cis-1(a) (OA)2, cis-2@(OA)2 and cis-3@(OA)2 was prepared by adding 5 μL of 60 mM respective standard solutions to 0.6 mL of 1 mM of OA in 10 mM of borate buffer solution. To this 4  $\mu L$ of 15 mM BMAN solution was added and the solution was purged with nitrogen for 20 min. The solution was irradiated using a filter (Corning CS 3-75, cut off wavelength 375 nm) and monitored over a period of time by recording <sup>1</sup>H NMR spectra. For energy transfer experiments, a solution containing cis-1@ (OA)2, cis-2@(OA)2 and cis-3@(OA)2 was prepared by adding 5  $\mu L$  of 60 mM respective standard solutions to 0.6 mL of 1 mM of OA in 10 mM of borate buffer solution. To this, 60 µL

of 20 mM 4-TMABP solution was added and the solution was irradiated using a filter (Corning CS 0-51, cut off wavelength 360 nm) and monitored over a period of time by recording <sup>1</sup>H NMR spectra. Experiments with corresponding *trans*-isomers were also repeated by following a similar procedure.

## Results

We have earlier reported inclusion of stilbenes within OA and their photochemistry in the confined environment of the OA capsule.39-43 This study concerns energy and electron transfer sensitized isomerization of OA encapsulated stilbenes. Although occurrence of electron transfer across the OA capsular wall has been established through ultrafast photophysical studies, 23,25-27 no electron transfer sensitized photoreaction of OA encapsulated molecules has been reported. The lack of

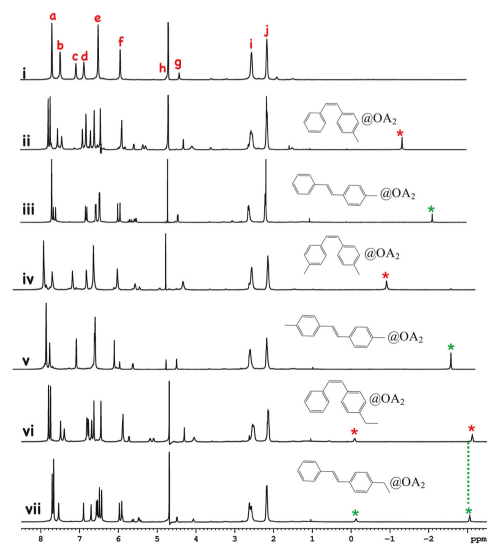


Fig. 1 <sup>1</sup>H NMR spectra (500 MHz, D<sub>2</sub>O) of (i) 1 mM OA in borate buffer, (ii) cis-1@(OA)<sub>2</sub>, (iii) trans-1@(OA)<sub>2</sub>, (iv) cis-2@(OA)<sub>2</sub>, (v) trans-2@(OA)<sub>2</sub>, (vi) cis-3@(OA)<sub>2</sub> and (vii) trans-3@(OA)<sub>2</sub> (\* & \*) – represent the bound methyl protons of (\*) cis and (\*) trans respectively. The OA signals are identified as alphabets; see Scheme 1 for the structure of OA and the identified protons.

clear understanding of the location of the sensitizer in the known examples of triplet sensitized isomerization of encapsulated olefins prompted us to reexamine this phenomenon.  $^{40,41}$  To keep the capsule and the sensitizers closer, cationic molecules were chosen as sensitizers so that they would be electrostatically held closer to the encapsulated stilbenes by the anionic OA. To infer the location of the sensitizers,  $^1\mathrm{H}$  NMR spectra of stilbene@OA2 capsule in the absence and presence of sensitizers were recorded. Initial experiments were focused on confirming the inclusion of stilbenes within OA and obtaining information on the guest–host ratio.

In Fig. 1, the <sup>1</sup>H NMR spectra of the two isomers of stilbene 1–3 in the presence of OA (1:2, guest to host ratio) are displayed. Inclusion of the guest within OA is evident from the large upfield shift of the 4-alkyl group. <sup>41</sup> Such an upfield shift is established to be a characteristic of the guest present in the cavity of OA that provides diamagnetic shielding. <sup>44–46</sup> Focusing on the alkyl signals, it is clear that in all three cases the signals for the *cis* and *trans* isomers appear with distinctly different chemical shifts. In the case of 3 the signals for the two isomers although closer are still distinct. This feature enabled us to follow the progress of irradiation by <sup>1</sup>H NMR.

Based on our earlier studies we are aware that stilbenes 1-3 form 1:2 guest-host complexes.<sup>39,40</sup> Formation of a capsular complex was further confirmed by measuring its diffusion constant by DOSY experiments. Values of diffusion constants are

summarized in the ESI (Table S1†). The values obtained for various complexes lower than that for free OA and the sensitizers are consistent with the numbers reported in the literature for the 1:2 (guest-host) complex.<sup>44,46</sup>

<sup>1</sup>H NMR spectra of cis-1@(OA)<sub>2</sub> in the presence of BMAN and trans-1@(OA)2 in the presence of 4-TMABP are shown in Fig. 2 and 3. Interestingly, the signals due to the guest methyl group and select hydrogens of OA are upfield shifted upon addition of the cationic sensitizers (see the green and red stars in the figure). The expanded spectrum included in each figure clearly reveals the upfield shift of the methyl signal with the increasing concentration of the cationic sensitizers. The fact that the sensitizer has a significant effect on the chemical shifts of the guest and host signals suggests that it remains closer to the host molecule. The lack of shift in the signals of the sensitizer in the presence of OA suggests that the sensitizer is not included within the capsule and it is present outside in water. Based on the effect of the sensitizer on the guest chemical shift we believe that the sensitizer is associated with the OA capsule as shown in Fig. 4. This is also confirmed by the diffusion constant of the sensitizers in the presence of the capsule. For example while BMAN and TMABP have diffusion constants of 4.35 and  $5.50 \times 10^{-6}$  cm<sup>2</sup> s<sup>-1</sup>, the termolecular complexes (stilbene@OA/sensitizer) have 1.29 and 1.88 × 10<sup>-6</sup> cm<sup>2</sup> s<sup>-1</sup> respectively (Table S1 in the ESI†). Decreased diffusion constants must be the result of the sensiztizers being attached

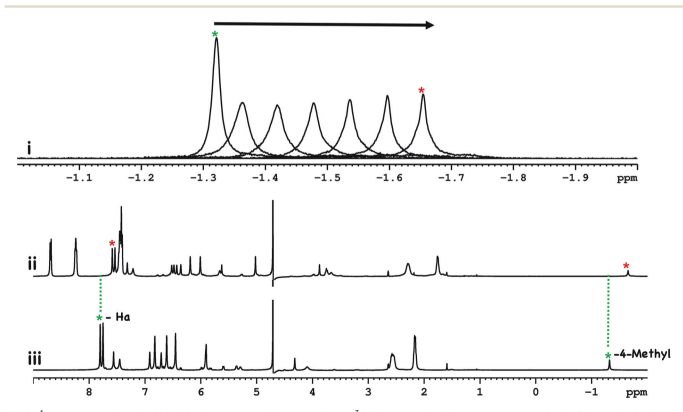


Fig. 2  $^{1}$ H NMR spectra (500 MHz) of *cis*-1 encapsulated within OA (1 × 10<sup>-3</sup> M) in buffered D<sub>2</sub>O in the presence of BMAN. (i) NMR signal of methyl protons of *cis*-1@(OA)<sub>2</sub> upon slow addition of BMAN (0.33 mM to 2.0 mM) and (ii) full spectrum upon addition of 2.0 mM of BMAN to *cis*-1@(OA)<sub>2</sub>. (iii) Full spectrum of *cis*-1@(OA)<sub>2</sub>; no BMAN. In the above spectra ( $\frac{1}{2}$ ) – indicates the signals in the absence of BMAN and ( $\frac{1}{2}$ ) – indicates the signals in the presence of BMAN.

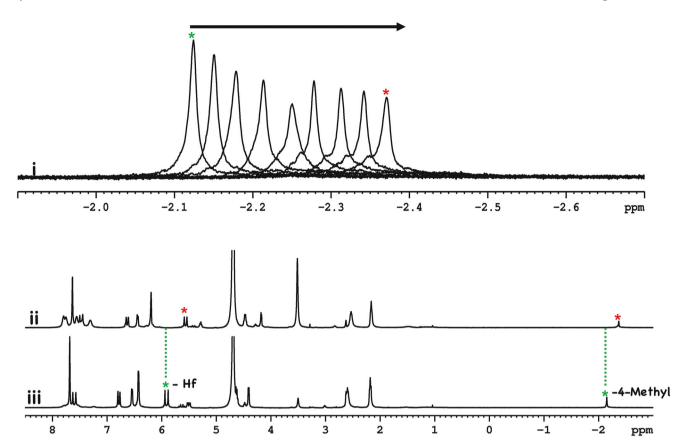


Fig. 3  $^{1}$ H NMR spectra (500 MHz) of *trans-*1 encapsulated within OA (1  $\times$  10<sup>-3</sup> M) in buffered D<sub>2</sub>O in the presence of 4-TMABP. (i) NMR signal of methyl protons of *trans-*1@(OA)<sub>2</sub> upon slow addition of 4-TMABP (0.33 mM to 2.0 mM). (ii) Full spectrum upon addition of 2.0 mM of 4-TMABP to *trans-*1@(OA)<sub>2</sub>. (iii) Full spectrum of *trans-*1@(OA)<sub>2</sub>; no 4-TMABP. In the above spectra (\*) – indicates the signals in the absence of 4-TMABP and (\*) – indicates the signals in the presence of 4-TMABP.

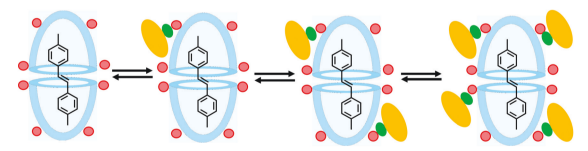


Fig. 4 Graphical representation of association of the cationic sensitizer to the capsule. We believe that with increasing concentration of the sensitizer the number of sensitizer molecules attached to the capsule would increase. Red circles represent COO<sup>-</sup> and green ovals (CH<sub>3</sub>)<sub>3</sub>N<sup>+</sup>.

to the capsules. The number of molecules attached to the capsule is expected to increase with the concentration which is likely to be responsible for the chemical shift dependence on the sensitizer concentration. The upfield shift in the presence of cationic sensitizers, most likely, is the result of changes in the electron density of the aromatics that form the capsular wall.

Having established the nature of the complexes and location of the sensitizers we proceeded to irradiate the

stilbene@OA<sub>2</sub>-sensitizer complexes in borate buffer. Control experiments showed that these solutions were stable in the dark for days. Since host OA is known to act as an energy and electron transfer sensitizer<sup>47</sup> and stilbenes photoisomerize upon direct light absorption<sup>35</sup> we made sure that light is absorbed only by the sensitizer. The absorption spectra of the complex, sensitizers and the filters used are provided in Fig. 5. Under our irradiation conditions (with Corning filters 0-51 and 3-75) the light is absorbed mainly by the sensitizers.

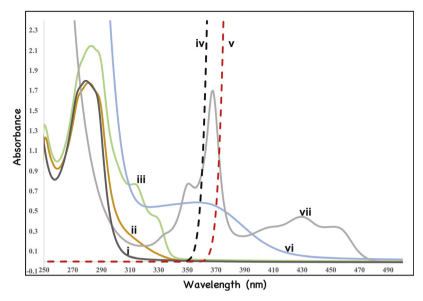


Fig. 5 Absorption spectra of (i) OA ([OA] =  $1 \times 10^{-4}$  M), (ii) cis-1@(OA)<sub>2</sub> ([cis-1] =  $1 \times 10^{-4}$  M: [OA] =  $2 \times 10^{-4}$  M), (iii) trans-1@(OA)<sub>2</sub> ([trans-1] =  $1 \times 10^{-4}$  M: [OA] =  $2 \times 10^{-4}$  M), (iv) filter 0-51, (v) filter 3-75, (vi) 4-TMABP ([4-TMABP] =  $0.5 \times 10^{-4}$  M) and (vii) BMAN ([BMAN] =  $0.5 \times 10^{-4}$  M).

Therefore, geometric isomerization reported here we believe is mainly due to the sensitization by BMAN and 4-TMABP and not due to direct absorption by encapsulated stilbenes. To confirm this, we carried out control experiments for the stilbene complexes using the same filter but without sensitizers. Less than 15% isomerization occurred under these conditions. Results of this irradiation are included in Table 1.

We chose BMAN to test the feasibility of electron transfer sensitized isomerization across the OA wall. Sensitization with BMAN was carried out using a Corning 3-75 filter (Fig. 5). Prior to this study we attempted sensitization with dimethyl viologen, N-methylacridinium iodide, dimethyldiazaphenanthrenium iodide and dimethyldiazapyrenium iodide. 47 But none of them were effective in generating the cis radical cation. Various problems confounded the choice: absorption in the case of dimethyl viologen, poor stability in water in the case of N-methylacridinium iodide and not enough oxidizing power in the case of the last two. Therefore, BMAN with stronger oxidizing power and longer absorption seemed promising.  $^{48-50}$ As seen in Table 1 it was effective as the sensitizer. Photoisomerization occurred in the time range of 4–10 h. <sup>1</sup>H NMR spectra of the irradiated samples of cis-1@(OA)2 are shown in Fig. 6. The spectra for others are provided in the ESI (Fig. S20 and S21†). Clearly, the cis isomer gave the trans without any side reactions and as expected conversion was 100%. However, BMAN was found to be unstable in water upon prolonged (>5 h) irradiation.<sup>50</sup> But the amount of BMAN used was sufficient to function as the sensitizer.

In the case of 4-TMABP sensitization, six samples (*cis* and *trans* isomers of 1, 2 and 3) along with controls were irradiated using light from a 450 W medium pressure mercury lamp filtered through a Corning 0-51 filter (Fig. 5). The concentrations of the stilbenes (1, 2 and 3) and 4-TMABP were 0.5 mM and

 Table 1
 Product distribution upon energy and electron transfer sensitized geometric isomerization of OA encapsulated stilbenes

Photostationary state during energy transfer sensitization with 4-TMABP <sup>a,b,c</sup>				Product distribution during electron transfer sensitization by BMAN <sup>c,d,e</sup>	
Starting isomer: 100% cis		Starting isomer: 100% trans		Starting isomer:100% cis	
cis	trans	cis	trans	cis	trans
50	50	49	51	10	90
85	15	5	95	90	10
0	100	0	100	0	100
81	19	0	100	85	15
84	16	85	15	73	27
96	4	14	86	100	0
	Star ison 100 cis 50 85 0 81 84	during ener sensitization 4-TMABP <sup>a,b,c</sup> Starting isomer: 100% cis  cis trans  50 50 85 15 0 100 81 19 84 16	during energy trasensitization with 4-TMABP <sup>a,b,c</sup> Starting isor isomer: 100° cis transcis trans cis  50 50 49 85 15 5 0 100 0 81 19 0 84 16 85	during energy transfer sensitization with 4-TMABP <sup>a,b,c</sup> Starting isomer: 100% 100% cis trans           50         50         49         51           85         15         5         95           0         100         0         100           81         19         0         100           84         16         85         15	during energy transfer sensitization with 4-TMABP <sup>a,b,c</sup>   Starting   Starting   isomer:   isomer:   100%   cis   trans   isomer: 100%   cis   trans   cis   trans   cis   cis   trans   cis   cis   trans   cis   trans   cis   cis   trans   cis   cis   trans   cis   cis

 $^a$  Irradiation times for energy transfer experiments were  $\sim$ 72 h; numbers entered are an average of 3 runs.  $^b$  Note that irradiation was performed starting from pure cis and pure trans.  $^c$  Control experiments did not use sensitizers.  $^d$  Irradiation times for electron transfer experiments varied between 4 and 10 h; numbers entered are an average of 3 runs.  $^e$  Note that irradiation was performed starting from pure cis only.

2 mM, respectively. The control without 4-TMABP under the same conditions was carried out. The progress of the reaction was followed by recording <sup>1</sup>H NMR and integrating the signals due to the 4-alkyl group of the stilbenes. Because of the use of cut-off filters it took >70 h to reach the photostationary state (pss). The percentage of *trans* and *cis* isomers at the pss starting from *trans* and *cis* isomers is provided in Table 1. <sup>1</sup>H NMR spectra of 4-TMABP sensitized isomerization of 1@(OA)<sub>2</sub> are shown in Fig. 7. For the other stilbenes the spectra are pro-

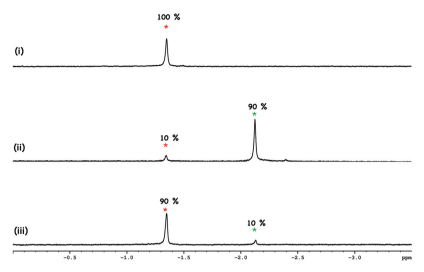


Fig. 6 Partial  $^{1}$ H NMR spectra (500 MHz) of cis-1@(OA)<sub>2</sub> encapsulated within OA (1 × 10<sup>-3</sup> M) in buffered D<sub>2</sub>O after addition of photocatalyst BMAN (i) before irradiation, (ii) after 4 hours irradiation and (iii) without BMAN after 24 hours irradiation (control).

vided in the ESI (Fig. S18 and S19†). It is important to note that isomerization was much less effective in the absence of the sensitizer (control). For example, in the case of 1@(OA)<sub>2</sub>, in the absence of the sensitizer the *cis* isomerized to 15% *trans* while upon sensitization it gave 50% *trans*. Similarly, the *trans*-1 gave 5% *cis* by direct light absorption and 49% by sensitization. Similar observations were made with other two stilbenes 2 and 3. These observations clearly establish that 4-TMABP functions as a sensitizer in the photoisomerizations listed in Table 1. The fact that both *cis* and *trans* isomers could be sensitized suggests certainly that sensitization is not through electron transfer but energy transfer.

## Discussion

Photoinduced electron transfer sensitized geometric isomerization of stilbenes has been investigated in detail by the Lewis group. <sup>36,37</sup> Sensitization by 9-cyanoanthracene, 9,10-dicyanoanthracene, 2,6,9,10-tetracyanoanthracene and *N*-methylacridinium hexafluorophosphate (NMA) is demonstrated to be effective in converting *cis*-stilbenes to the corresponding *trans* isomers. First three are not water soluble and too large to fit within an OA capsule. Although *N*-methylacridinium salt is water soluble, prolonged irradiation resulted in its degradation. To overcome this problem we employed 9-mesityl *N*-methylacridinium salt. <sup>51</sup> Similar to NMA this also had poor stability in water. In addition to their poor stability the two were found to be ineffective in sensitizing the isomerization of the stilbenes.

Having eliminated the common electron transfer sensitizers, we used the less well known sensitizer BMAN (dimeric form of NMA) for inducing geometric isomerization of the encapsulated *cis*-stilbenes.<sup>50</sup> Its absorption in the visible region extending up to 480 nm (2.58 eV) permitted excitation

of the sensitizer without directly exciting the stilbenes. Also its reduction potential is reported to be -0.30 eV in water where the sensitizer resides under our condition. Based on the Rehm-Weller equation the free energy of electron transfer from excited BMAN to cis-stilbene is estimated to be exothermic by ~0.2 eV. For the methyl substituted stilbenes the process is even more exothermic than that for unsubstituted stilbenes. Given that the OA has a lower oxidation potential than cis-stilbene, it could also act as an electron donor to the excited BMAN. Based on the estimated oxidation potential of OA (~1.5 eV) the free energy of electron transfer from excited BMAN to OA is estimated to be exothermic by ~0.7 eV. As shown in Fig. 8, all three, host, guest and host-guest complex namely OA, cis-1 and cis-1@OA2 quench the fluorescence of BMAN. Since OA as well as cis-1 can act as quenchers, in the case of cis-1 @OA2 it is not obvious what is the quencher, OA or cis-1. We believe that both are acting as quenchers under our conditions. Since the electron transfer from cis-stilbene to the OA radical cation is an uphill process (+~0.5 eV; oxidation potential of cis-stilbene and OA is 2.06 eV and ~1.5 eV, respectively), we believe any reaction that we observe must be the result of direct quenching of the excited BMAN by encapsulated cis-stilbene.

Irradiation (>370 nm; 450 W medium pressure mercury lamp with Corning CS 3-75) of a solution of OA encapsulated *cis*-stilbene and BMAN (0.5 mM and 0.1 mM) gave the corresponding *trans* isomer quantitatively within 4 h-10 h. Under the same conditions without BMAN the isomerization was slow and 10% *trans* isomer was formed even after 24 h of irradiation. As shown in Fig. 6 and Fig. S20, S21† for the three stilbenes (1-3) the isomerization is clean and no other products were formed. We believe the results support the conclusion that electron transfer from stilbenes to excited BMAN does occur across the walls of the OA capsule. Clearly, the photoreaction of encapsulated molecules can be initiated by

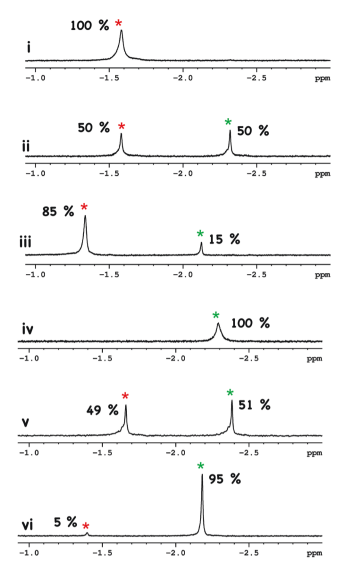


Fig. 7 Partial <sup>1</sup>H NMR spectra (500 MHz) of cis-1@(OA)<sub>2</sub> and trans-1@ (OA)<sub>2</sub> encapsulated within OA (1 × 10<sup>-3</sup> M) in buffered D<sub>2</sub>O after addition of sensitizer 4-TMABP (i & iv) initial, (ii & v) after 72 hours irradiation and (iii & vi) without 4-TMABP after 72 hours irradiation (control). ( $\frac{1}{2}$  &  $\frac{1}{2}$ ) represent the bound methyl protons of ( $\frac{1}{2}$ ) cis and ( $\frac{1}{2}$ ) trans, respectively.

molecules that stay outside the capsule. Such a possibility opens an opportunity to remote trigger photoreactions *via* electron transfer. While the results presented here establish the phenomenon, we are yet to find a good electron transfer sensitizer that would be stable and function in water. Unfortunately, BMAN degrades slowly in water under our irradiation conditions. This prompts us to continue our search for a better electron transfer sensitizer in the context of remote sensitization in water.

We visualized that a combination of supramolecular photochemistry<sup>11,19,52</sup> and triplet-triplet energy transfer<sup>4</sup> would be a valuable tool to manipulate excited state reactions. Following the remarkable discovery of triplet-triplet energy transfer in a rigid solution at 77 K by Terenin and Ermolaev,<sup>53</sup> the value of this process in organic photochemistry was estab-

lished by Hammond and his students in the 1960s. 35,54-59 Recently, this technique has re-emerged in the context of finding new green synthetic methods to build organic molecules. 7,10 To be effective as the triplet sensitizer under our conditions, the sensitizer must remain closer to the capsule and also should be water-soluble. However, the common wellknown organic sensitizers are generally water insoluble. We overcame this problem by functionalizing the well-known sensitizer benzophenone with an ammonium group that would not only make them water soluble but also get them non-covalently attached to the anionic capsule. As evident from the <sup>1</sup>H NMR spectra in Fig. 3, sensitizer 4-TMABP remains closer to the capsule in aqueous solution. The up-field shift of the <sup>1</sup>H signals of the host as well as the guest with the increasing concentration of the sensitizer is a direct result of the proximity of the capsule and the sensitizer. This feature we believe will facilitate double electron transfer (collisional energy transfer) across the capsular wall.

The emission spectrum of TMABP in aqueous solution at room temperature is shown in Fig. S17 in the ESI.† Since the emission is not quenched by oxygen we believe this to be fluorescence. Based on the 0-0 band the S<sub>1</sub> energy of the sensitizer is estimated to be 71 kcal mol<sup>-1</sup> (400 nm). Currently, we do not know the nature of the lowest excited state ( $n\pi^*$  or  $\pi\pi^*$ ). Independent of this, it is safe to assume the lowest triplet of 4-TMABP to be above 60 kcal mol<sup>-1</sup> (benzophenone: 69 kcal mol<sup>-1</sup>). With the triplet energies of *cis* and *trans*-stilbenes estimated to be 57 and 49 kcal mol<sup>-1</sup> respectively, we believe that 4-TMABP would be able to sensitize both isomers. The three stilbenes chosen for investigation have unique behavior within OA. For example, within the OA capsule the three chosen olefins behave differently:<sup>42</sup> at the pss cis and trans isomers are present in equal amounts in 1, trans rich in the case of 2, and cis rich in the case of 3. Therefore, based on the photostationary state achieved one could infer whether the reaction occurred within OA or in aqueous solution. With this background we analyze the results summarized in Table 1.

Irradiation was performed with light of wavelength >350 nm where the stilbenes do not absorb. Progress of isomerization was followed by <sup>1</sup>H NMR (Fig. 7 and Fig. S18, S19 in the ESI†). Independent of the initial isomer, the same photostationary state (pss) was achieved in about 72 h. The control experiments revealed that although there is background isomerization, the majority of the isomerization observed in the presence of 4-TMABP is the result of sensitization. Considering the absorptions of stilbene and 4-TMABP (Fig. 5), one would expect 4-TMABP to be the primary absorber under our irradiation conditions. Close examination of the table reveals that for example when 1@(OA)2 was irradiated in the absence of the sensitizer the pss contained 5% cis and 95% trans when the starting isomer was trans, but the pss was 85% cis and 15% trans when the starting isomer was cis. Clearly the pss ratio is not the same starting from the two isomers. On the other hand, when [1@(OA)<sub>2</sub> + 4-TMABP] solution was irradiated the same cis-trans composition was obtained at the pss (equal amounts of *cis* and *trans*). If the iso-

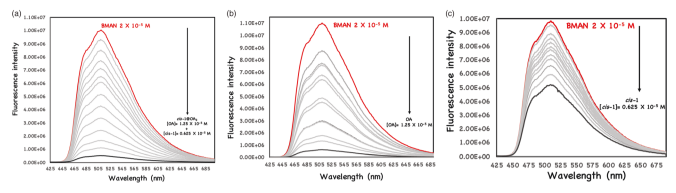


Fig. 8 (a) Emission spectra ( $\lambda_{exc}$  = 420 nm) of acceptor BMAN (red), and BMAN (black) as a function of the donor (cis-1@OA<sub>2</sub>) concentration;  $([BMAN] = 2.0 \times 10^{-5} \text{ M}, [cis-1] = 0 - 0.625 \times 10^{-5} \text{ M} + [OA] = 0 - 1.25 \times 10^{-5} \text{ M}$  in buffer/H<sub>2</sub>O, pH = 8.7). (b) Emission spectra ( $\lambda_{exc} = 420 \text{ nm}$ ) of acceptance tor BMAN (red) and BMAN (black) as a function of the donor OA concentration; ([BMAN] =  $2.0 \times 10^{-5}$  M, [OA] =  $0-1.25 \times 10^{-5}$  M in buffer/H<sub>2</sub>O, pH = 8.7). (c) Emission spectra ( $\lambda_{exc}$  = 420 nm) of acceptor BMAN (red), and BMAN (black) as a function of the donor cis-1 concentration; ([BMAN] = 2.0 ×  $10^{-5}$  M, [cis-1] =  $0-0.625 \times 10^{-5}$  M in buffer/H<sub>2</sub>O, pH = 8.7).

merization is due to direct absorption the ratio would not be the same from both isomers. More importantly, in the case of 2@(OA)<sub>2</sub> upon direct excitation about 19% of trans was obtained from cis. On the other hand, irradiation of [2@(OA)2 + 4-TMABP], 100% trans resulted at the pss. Once again this must be the result of sensitization by 4-TMABP, not due to direct excitation. Similar analysis of the results of 3@(OA)2 supports the conclusion that 4-TMABP is able to generate the triplet of encapsulated stilbenes from staying outside the capsule. In conclusion, results summarized in Table 1 suggest that the triplet-triplet energy transfer could occur across the OA wall.

### Conclusions

Results presented in this study demonstrate that sensitizers remaining outside a supramolecular assembly can trigger photoreactions of guest molecules trapped within the assembly. Based on ultrafast time resolved studies we are aware that in the time scale of excited states, the OA capsule remains closed.<sup>60</sup> Although at longer time (300 ms) scales disassembly-assembly is likely,61-64 in the time scale of excited states (µs to ns) the capsule is closed and the energy and electron transfer occurs across the molecular wall of the OA capsule. Thus the proof of principle of the viability of remote sensitization is established and this, in our opinion offers an opportunity to remotely manipulate photoreactions in a tight capsular assembly. We are in the process of identifying useful and stable sensitizers and reactions to explore the concept further.

### Conflicts of interest

There are no conflicts of interest to declare.

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