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Demonstration of an azobenzene derivative based solar thermal energy storage system[†]

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Molecules capable of reversible storage of solar energy have recently attracted increasing interest, and are often referred to as molecular solar thermal energy storage (MOST) systems. Azobenzene derivatives have great potential as an active MOST candidate. However, an operating lab scale experiment including solar energy capture/storage and release has still not been demonstrated. In the present work, a liquid azobenzene derivative is tested comprehensively for this purpose. The system features several attractive properties, such as a long energy storage half-life (40 h) at room temperature, as well as an excellent robustness demonstrated by optically charging and discharging the molecule over 203 cycles without any sign of degradation (total operation time of 23 h). Successful measurements of solar energy storage under simulated sunlight in a microfluidic chip device have been achieved. The identification of two heterogeneous catalyst systems during testing enabled the construction of a fixed bed flow reactor demonstrating catalyzed back-conversion from cis to trans azobenzene at room temperature under flow conditions. The working mechanism of the more suitable catalytic candidate was rationalized by detailed density functional theory (DFT) calculations. Thus, this work provides detailed insights into the azobenzene based MOST candidate and identifies where the system has to be improved for future solar energy storage applications.

Introduction

The development of sustainable energy has been attracting increasing attention, due to the pressing environmental and social challenges linked to high dependency on fossil fuels in our modern society.1 As the most abundant energy source for Earth, the Sun can provide the energy needed for mankind for an entire year in only six hours.2 Various ways to take advantage of solar energy have been studied and developed over the last few decades, including photovoltaics,3 artificial photosynthesis,4 and solar thermal heating.5 In order to store sunlight for future use, molecular solar thermal storage techniques (MOST, 6,7 also known as solar thermal fuels, STF8) focus on harvesting solar energy and storing it in photoswitchable materials. A parent molecule can be isomerized by solar irradiation to a metastable high energy photoisomer for a long storage period. When the energy is required, it can be thermally or catalytically back-converted to the parent state. Ideally, the stored energy should be released on demand as heat, thus operating as a closed-cycle system.

In order to realize the MOST concept, several features concerning the charge and discharge process need to be considered.^{7,9,10} (1) Since more than 50% of sunlight is distributed from 300 nm to 800 nm, the parent molecules should be able to absorb broadly in this spectral region. (2) The quantum yield of the photoisomerisation reaction should be as close to unity as possible. (3) The storage half-life at room temperature should be long enough to fulfill application-relevant storage times, such as daily, monthly, yearly or even longer storage requirements. (4) The energy of the metastable photoisomer should be significantly higher than the ground state of the parent isomer. (5) The system should ideally be able to operate over an infinite number of charging and heat releasing cycles. (6) Using a heterogeneous catalyst, the photoisomer should easily release the stored energy as heat. (7) For energy collection, storage, and bulk heating applications, MOST materials need to be pumped between solar collector, storage reservoir and heat extraction devices. As a consequence, the MOST system should be a liquid or highly soluble solid.

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The MOST concept has been represented by many potential candidates, including norbornadiene/quadricyclane derivatives, 7,9-14 dihydroazulene/vinylheptafulvene couples, 15-21 difulvalenediruthenium complexes,6 anthracene dimers,22 Dewar isomers²³ and azobenzene derivatives.^{8,24-43} The last systems have been in the spotlight recently due to the broad absorption spectrum, high robustness, daily storage half-life, tunable energy density (I mol⁻¹) of photoswitches, and the low synthetic cost.40-42 AZO1 (see Fig. 1a) possesses many of the needed features described above; however it has not been tested in a full MOST operating cycle including energy capture, storage and release. Here, for the first time, we investigate the performance of AZO1 by studying its function through multiple energy storage and release cycles. In order to do so, we have identified and tested a heterogeneous catalyst system for the energy release step and constructed a flow system consisting of a solar collector and a fixed-bed catalytic converter. Furthermore, the functioning mechanism of the catalytic conversion is further rationalized using proposed and PCM-B3LYP-D3BJ/6-31G(d)+SDD(Cu) calculations.

Results and discussion

To characterize the physical properties of AZO1, it was first studied in toluene in terms of its absorptivity, half-life of energy storage and quantum yields of the photoisomerization reaction. Comparing these results with available data of AZO1 in methanol,43 the maximum in the absorption spectrum was observed to be slightly red shifted, indicating a low solvent polarity effect compared to, for instance, a DHA-MOST system¹⁸ (ca. 5 nm,

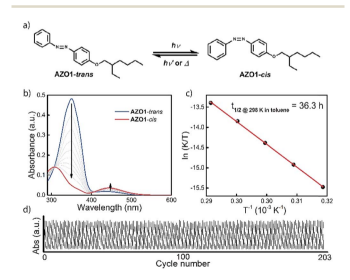


Fig. 1 (a) Structure of AZO1 in the trans and cis state. (b) Absorption spectra of AZO1-trans (in blue) and its corresponding photoisomer AZO1-cis (in red). The sample (ca. 2 mg in 100 mL toluene) was converted with a 340 nm LED lamp. (c) Eyring plot of AZO1-cis. The half-life at 25 °C was determined as 36.3 h in toluene solution. (d) Optical cycling test of AZO1 in toluene. The figure shows 203 cycles in total. The absorbance was recorded at 350 nm. The charge and discharge processes were archived with a 340 nm LED (\sim 60 mW) over 100 s and a 455 nm LED (~1020 mW) over 300 s, alternatively.

 $\varepsilon_{\text{Max} = 350 \text{ nm}} = 2.6 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$, see Fig. 1b). Concerning the storage lifetime, a half-life from cis to trans state of 36.3 h in toluene can be extracted from the Eyring plot. The activation enthalpy was calculated as $\Delta H_{\text{therm}}^{\ddagger} = 93.7 \text{ kJ mol}^{-1}$, together with an activation entropy $\Delta S_{\text{therm}}^{\ddagger} = -31.7 \text{ J mol}^{-1} \text{ K}^{-1}$ (S2, ESI†). The quantum yield of the trans to cis photoisomerisation reaction was determined to be 21% at 340 nm, using a literature procedure44 (S3, ESI†). In addition, it was observed that the AZO1-cis state can photoisomerise back to the trans form. The corresponding quantum yield of the cis to trans photoisomerisation was determined to be 23% at 455 nm, slightly higher than the reverse process. Due to this photo-induced twoway switch effect for AZO1, full conversion via full spectrum solar light can likely not be achieved and a band pass filter was used in the device conversion experiments.

Good cyclability is one of the most important criteria of MOST systems. To investigate the robustness of AZO1, a solution of ca. 10^{-5} M in toluene was prepared without degassing. Two controllable LED light sources were turned on and off alternatingly (340 nm with 100 s irradiation time and 455 nm with 300 s irradiation time) to charge and discharge the molecules back and forth. After 203 complete cycles (with a total operation time of 22.6 h), no significant signs of degradation were observed, thus demonstrating high robustness of AZO1, even in the presence of oxygen (see Fig. 1c).

To further demonstrate the functionality of the AZO1 system in a real device, a continuous flow system was built. For this labscale conversion test under an AM1.5 solar simulator, diluted solutions of AZO1 were used to collect data by UV-vis spectroscopy. Two concentrations of AZO1 solutions were pumped individually through a microfluidic chip,20,21 with varying residence time (33.9 mm³ inner volume with a channel depth of 100 μm, see Fig. 2a). Since AZO1-cis can be back-converted by visible light, an optical UV transmitting band pass filter (<400 nm) was inserted in between the solar simulator and the microfluidic chip. The UV-Vis spectra of the AZO1 solution before and after pumping into the microfluidic chip were recorded. Eqn (1) was used to calculate the cis-to-trans conversion percentage (S4, ESI†):

$$\text{Conversion \%} = \frac{\frac{A_{@350 \text{ nm}}}{A_{\text{iso}@306 \text{ nm}}} \varepsilon_{\text{iso}@306 \text{ nm}} - \varepsilon_{\text{isomer}@350 \text{ nm}}}{\varepsilon_{\text{parent}@350 \text{ nm}} - \varepsilon_{\text{isomer}@350 \text{ nm}}}$$
 (1)

where $A_{@350 \text{ nm}}$ is the actual absorbance at 350 nm; $A_{\text{iso}@306 \text{ nm}}$ and $\varepsilon_{iso@306 \text{ nm}}$ correspond to the absorbance and absorptivity of the solution at its isosbestic point at 306 nm, respectively; and $\varepsilon_{parent@350 \text{ nm}}$ and $\varepsilon_{isomer@350 \text{ nm}}$ are the absorptivity of the AZO1-trans and AZO1-cis isomers at 350 nm, respectively. A maximum conversion of around 80% from the AZO1 trans to cis state was obtained from a solution of 2×10^{-4} M. This is likely due to the photo-stationary state of azobenzene trans-to-cis photoisomerization (see Fig. 2b). Concerning the energy storage efficiency, 0.88% of the solar energy could theoretically be stored in a neat sample (S5, ESI†). The highest efficiency that can be expected for a 5×10^{-4} M solution is 0.02%, and for a 2 \times 10⁻⁴ M solution is 0.01%. With a band pass filter (SCHOTT,

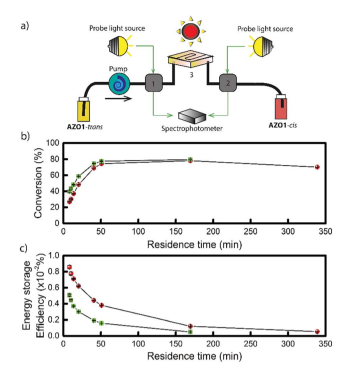


Fig. 2 (a) Experimental setup of AZO1 in a microfluidic chip device. Grav boxes 1 and 2 contain a flow UV-vis detection device connected to a portable spectrophotometer, 3, corresponding to a total volume of 33.9 mm³ quartz chip, with a 100 μ m optical path length. (b) Experimental data of the two concentrations used: 2×10^{-4} M in green and 5 \times 10 $^{-4}$ M in red. Conversion percentage of AZO1- transwith different residence times in the microfluidic chip. (c) Measured energy storage efficiency of the AZO1 compound in toluene.

UG11) and a solar simulator, the actual measurements showed that the maximum energy storage efficiency was ca. 0.009% for the 5 \times 10⁻⁴ M solution, and 0.005% for the 2 \times 10⁻⁴ M solution (see Fig. 2c, S6, ESI†). These experimental results were close to reach the theoretical predictions, but are still limited overall by the concentrations used and the photo-stationary state between the two species.

After solar capture/storage, energy release is the second fundamental process for the MOST concept. To estimate the adiabatic heat release as a function of concentration, a modified equation from a previous study11 was used (eqn (2)):

$$\Delta T = \frac{c M_{\rm w} \Delta H_{\rm storage}}{\frac{c^2 M_{\rm w}^2}{\rho_{\rm AZOI}} C_{\rm p_AZOI} + 1 - \frac{c M_{\rm w}}{\rho_{\rm AZOI}} \rho_{\rm solvent} C_{\rm p_solvent}}$$
(2)

where c is the concentration of AZO1, $M_{\rm w}$ represents its molecular weight; $\Delta H_{\text{storage}}$ corresponds to the DSC measured energy storage capacity of AZO1-cis which equals to 167.5 J g^{-1} ; ρ_{AZO1} is the volumetric mass density of AZO1 supposed to be 1.09 g mL⁻¹; C_{p_AZO1} is the specific heat capacity of AZO1 in J g⁻¹ K⁻¹, assumed similar to that of unsubstituted azobenzene;⁴⁵ and ρ_{solvent} and $C_{\text{p_solvent}}$ address the volumetric mass density in g L^{-1} and the specific heat capacity in J g^{-1} K^{-1} of the solvent, respectively (which are equal to 867 g L⁻¹ and 1.7 J g⁻¹ K⁻¹). In this approach, the volume load factor of the solvent and

photoisomer was considered, i.e. when the concentration $\left(1 - rac{cM_{
m w}}{
ho_{
m AZO1}}
ight)
ho_{
m solvent}C_{
m p_solvent}$ approaches conditions, approaches zero. This correction is reasonable for all MOST heat release estimations with close to neat conditions. In the case of AZO1, the theoretical maximum temperature difference was calculated as ca. 226 °C for a fully charged neat sample (see

Concerning related catalysts, several candidates, including mineral acids like perchloric acid, Cu(II) salts (CuCl₂,Cu(OAc)₂), gold nanoparticles which involve a redox mechanism, as well as electrocatalytic methods have been reported to induce the backconversion of azobenzene derivatives. 46-48 However, for a closed cycle system which can be operated in devices, a heterogeneous catalyst that can be fixed in a reaction centre is required. Based on this fact, a heterogeneous catalyst or an insoluble homogeneous catalyst needs to be developed. For AZO1, two potential catalysts, cobalt(II) phthalocyanine physisorbed on the surface of activated carbon (CoPc@C) and [Cu(CH₃CN)₄]PF₆ fulfill the described physical properties and thus, they were tested individually. Both showed a positive effect on reducing the backconversion half-life at room temperature. [Cu(CH₃CN)₄]PF₆ has a very low solubility in toluene, being active for various MOST systems, including norbornadiene/quadricyclane derivatives11 and dihydroazulene/vinylheptafulvene couples.20,49 For the AZO1-cis compound, a back-conversion reaction rate of up to 30 s⁻¹ was calculated at room temperature of 25 $^{\circ}$ C (up to 6 \times 10^6 times higher than a reaction rate of 5×10^{-6} s⁻¹ without the catalyst at 25 °C; see Fig. 3a, S7, ESI†). CoPc@C was produced following a reported procedure; however it had a risk of leaching from its solid support.11 Therefore, the [Cu(CH3CN)4]PF6 salt was chosen to be incorporated into the catalytic device for further testing. With this result in mind, a small-sized reaction centre was built. 5 mg of Cu(I) salt was inserted into a Teflon tube which has a 1 mm inner diameter (see Fig. 3b). 5×10^{-4} M 79% **AZO1**-cis solution from the microfluidic chip experiments was then flowed through the catalytic bed with a speed of 1 mL h⁻¹. As result, 48% of **AZO1**-cis was successfully back-converted to the corresponding trans state. Thus, the described continuous fluidic chip experiments can achieve the requirements for the complete application of the MOST concept including photon capture/storage and energy release processes.

To further understand the mechanism of back-conversion with $[Cu(CH_3CN)_4]PF_6$, a detailed study within the framework of the density functional theory (DFT) was performed. A simplified AZO1 system was used replacing the ethylhexyl moiety with a methyl group at the PCM-B3LYP-D3BJ/6-31G(d)+SDD(Cu) level of theory (S8, ESI†). Among the different coordination options, the CH₃CN ligands could be displaced by the new ligand (the cis isomer of the azobenzene moiety, in this case). Four different coordination possibilities were considered to ensure a good modelling of the experimental conditions. In all cases, the formation of the new complex is slightly endergonic (1-5 kcal mol⁻¹). As the azobenzene derivative is not symmetric, two sets of coordination alternatives arise. Coordination through the nitrogen atom directly linked to the

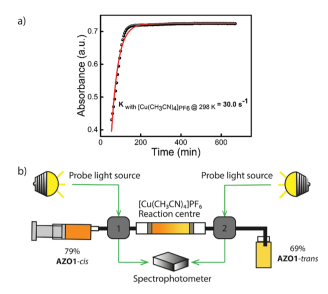


Fig. 3 (a) Kinetics study of AZO1-cis in the presence of [Cu(CH₃CN)₄] PF₆ in toluene at 25 °C. Black dots represent the experimental data, red curve corresponds to the exponential fitting. The reaction constant was calculated as 30.0 s⁻¹ (b) Conceptual device demonstration of catalytic back-conversion of AZO1-cis. Around 5 mg of Cu(I) salt were loaded in to the reaction centre. A 5 \times 10⁻⁴ M 79% AZO1-cis solution was flown through the Cu(1) reactor with a speed of 1 mL h^{-1} . 69% of the AZO1-trans was obtained after the reaction centre, i.e. 48% of the AZO1-cis can be successfully back-converted to AZO1-trans state.

unsubstituted phenyl ring (path b, S8, ESI† and Fig. 4a) is favored (by ca. 2.3 kcal mol⁻¹) as compared to the methoxyphenyl linked nitrogen atom (path a). Isomerization of azobenzenes is very dependent on substitution and reaction conditions. 50 In this case, different pathways were considered and rotation and inversion mechanisms of the isomerization were explicitly evaluated. As in the not catalyzed thermal reaction,47 the inversion transition states (TS1a-TS1b) were found to be preferred.

A relaxed scan along the rotation coordinate (CNNC dihedral) (see Fig. 4b) reveals a possible S₀-T₁-S₀ mechanism accessible by coordination of the nitrogen atom of AZO1-cis to copper, which reveals a MLCT (metal-to-ligand charge transfer). In turn, this implies the participation of a π^* orbital of the azobenzene allowing rotation along the N=N with participation of the T₁ state. At relative high torsion angles, the triplet state becomes less energetic and this path yields a barrier of around 20 kcal mol⁻¹. This alternative mechanism allows for the bypassing of the more energetic inversion TS in the ground state $(30 \text{ kcal mol}^{-1})$ and it should be available at room temperature.

Thus, copper coordination yields a decrease in the isomerization barrier of ca. 6 kcal mol⁻¹ (computed barrier decreases from 25.4 kcal mol⁻¹ for free azobenzene to 19.4 kcal mol⁻¹). This relatively small decrease in the energy barrier is enough to allow the isomerization process to occur in a few minutes instead of some days, thus accelerating the isomerization by 4 orders of magnitude, resulting in good agreement with experiments (energy difference of 8 kcal mol⁻¹, from 23.5 kcal mol⁻¹ for the free AZO1 to 15.6 kcal mol⁻¹ for the copper activated

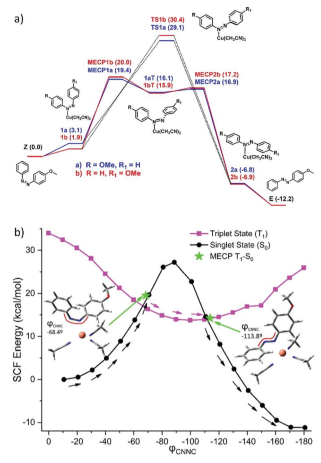


Fig. 4 (a) Reaction paths for cis-trans isomerization. Coordination through the nitrogen atom directly linked to the phenyl ring (path b, in red) and to the methoxyphenyl nitrogen atom (path a, in blue). Solid lines are the $S_0-T_1-S_0$ pathway (MECP1(a,b)-1(a,b)T-MECP2(a,b)). Dotted black lines are the singlet reaction path (TS1a-TS1b). Free energies referred to cis-AZO + Cu(CH₃CN)₄⁺. (b) Relaxed scan along CNNC rotation and MECPs

reaction). Overall, while [Cu(CH₃CN)₄]PF₆ is useful to increase the reaction rate notably, there is still great potential for improvement in the design of the energy release step. This could be done, for instance, favoring the MLCT or stabilizing the MECP between singlet and triplet states to increase the efficiency of the crossing. In both cases, this could lead to a decrease in the energy barrier and a subsequent acceleration of the back-conversion.

Conclusions

An AZO1-MOST charging/discharging cycle has been successfully demonstrated, including the study of the photophysical properties and back-conversion of the photoisomer. The AZO1 system features strong absorption ($\varepsilon_{\text{Max}@350 \text{ nm}} = 2.6 \times$ 10⁴ M⁻¹ cm⁻¹) and long thermal half-life (36.3 h). The optical cyclability test for a diluted solution shows no significant degradation in the presence of oxygen, allowing the use of AZO1 in toluene in application. For MOST application purposes, the conversion quantum yield at 340 nm was determined to be 21%

which is similar to the quantum yield of back-conversion of 23% at 455 nm. This implies that even diluted samples cannot be fully converted under sunlight. The estimated maximum energy storage efficiency for AZO1 in the neat state was calculated to be 0.88%. To experimentally demonstrate the functionality of this liquid azobenzene for MOST applications, conversion experiments were performed at different concentrations of AZO1 in a continuous microfluidic chip system. The measured maximum energy storage efficiency for a 5×10^{-4} M solution can reach close to 0.009%, close to reach the theoretical prediction of 0.02% at such a concentration. This, unfortunately, showed a strong photo-stationary effect between AZOcis and AZO-trans, therefore highlights the need to further develop the azobenzene system by altering the optical properties. 51-53 Further improvement to increase quantum yields and diminish the absorption of photoisomer, as well as differentiate the spectral difference between cis and trans states, would be certainly required. To estimate the theoretical heat release temperature difference, a corrected formula was introduced to account for the change from the diluted solution to the neat sample. A maximum temperature increase of 226 °C can be theoretically achieved by using eqn (2) with a correction term for a neat MOST sample. Furthermore, two new catalysts have been identified to allow the back-conversion of AZO1-cis and a reaction centre based on a Cu(1) salt has been prepared as a proof of concept for the energy release process. With a concentration of 5×10^{-4} M and a flow speed of 1 mL h⁻¹, 48% of the *cis* state isomer was successfully converted back to the trans state. However, the reaction rate was still too low to be used for macroscopic heat release purposes. Finally, a detailed theoretical study of the mechanism has been proposed to further understand the discharge process using a Cu(1) salt. This could help in the future design of new azobenzene derivatives and the corresponding catalysts. In addition, results shown here could be extended to the control of azobenzene derivatives for different applications.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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