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Evaluating Dihydroazulene/Vinylheptafulvene Photoswitches for Solar Energy Storage Applications

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Efficient solar energy storage is a key challenge in striving toward a sustainable future. For this reason, molecules capable of solar energy storage and release through valence isomerization, for so-called molecular solar thermal energy storage (MOST), have been investigated. Energy storage by photoconversion of the dihydroazulene/vinylheptafulvene (DHA/VHF) photothermal couple has been evaluated. The robust nature of this system is determined through multiple energy storage and release cycles at elevated temperatures in three different solvents. In a nonpolar solvent such as toluene, the DHA/VHF system can be cycled more than 70 times with less than 0.01% degradation per cycle. Moreover, the [Cu(CH₃CN)₄]PF₆-catalyzed conversion of VHF into DHA was demonstrated in a flow reactor. The performance of the DHA/VHF couple was also evaluated in prototype photoconversion devices, both in the laboratory by using a flow chip under simulated sunlight and under outdoor conditions by using a parabolic mirror. Device experiments demonstrated a solar energy storage efficiency of up to 0.13% in the chip device and up to 0.02% in the parabolic collector. Avenues for future improvements and optimization of the system are also discussed.

Introduction

The rise in energy demands as a consequence of fast economic growth and an increasing global population poses challenges for future generations. In 2050, energy consumption is projected to be 27.6 TW, twice that of 2001.^[1] At the same time, associated environmental challenges render the use of fossil fuels less desirable. Consequently, development of sustainable resources has become a focus of research during recent decades.^[2] As the most abundant energy source on Earth, solar energy utilization has seen a rapid expansion with more than 900% increase of installed capacity between 2009 and 2015, including both utility-scale solar power plants and distributed photovoltaics,^[3] as well as solar water heating for domestic and industrial applications.^[4] As solar energy production is being widely implemented, challenges associated with time variations in solar energy output and variations in energy consumption lead to an increased need for efficient, scalable solar energy storage technologies.

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One possible means for facilitating this need is through socalled molecular solar thermal energy storage (MOST) systems.^[5] A MOST system is based on a molecule that, upon exposure to sunlight, can be photoisomerized into a high-energy, metastable isomer. When required, the energy stored in the chemical bonds can be released by thermally or catalytically triggering back-conversion into the parent compound, thereby releasing the stored energy as heat. The requirements for such a system have been discussed in detail previously.^[6] In essence, an ideal MOST system should have a long lifetime under ambient conditions, high photoisomerization quantum yield, large spectral overlap with the solar spectrum, and sufficiently high energy of the metastable isomer. Furthermore, it should be highly robust and thus able to undergo multiple cycles without significant loss of capabilities through degradation.

Several types of photoswitches for MOST applications have been investigated, including functionalized norbornadienes,^[5a, 6a,b] fulvalene–diruthenium derivatives,^[2,7] azobenzenes,^[6c,8] azopolymers,^[9] and several dihydroazulene/vinylheptafulvene (DHA/VHF) derivatives.^[10] The latter system, which is shown in Scheme 1 (DHA1/VHF1), is the subject of the present work. DHA1 can be converted into its corresponding highenergy isomer VHF1 by a photoinduced ring-opening reaction. However, VHF can only undergo a thermal back reaction to the DHA form. Therefore, it should be possible to have full conversion between the two states in both directions. VHF1 itself exists in an equilibrium between s-cis and s-trans conformers; the s-trans conformer usually being the more stable of the two, whereas s-cis is the conformation necessary for ring closure. The combination of a large spectral difference between DHA and VHF and a high enthalpy of the photoreaction



Scheme 1. The dihydroazulene (DHA1)/vinylheptafulvene (VHF1)-based molecular solar thermal energy storage (MOST) system examined in this study.

of 35.2 kJ mol⁻¹ (correspond to 0.14 MJ kg⁻¹ calculated in vacuum, and 0.13 J cm⁻³ assuming a volumetric mass density of DHA of 0.9 g cm⁻³)^[10b] fulfills several of the requirements preferred for a MOST system.^[5a] In addition, many DHA derivatives have been made, where the lifetimes can be prolonged for the corresponding VHFs.^[11] Moreover, the synthesis of **DHA1** employs readily available starting materials and has been accomplished on a large (15 g) scale.^[12]

Herein, we investigate the DHA/VHF couple as a possible candidate in a MOST device. As a part of this study, we have assessed both solubility and stability of this system by subjecting DHA/VHF to multiple photothermal energy storage and release cycles. In addition, we test the conversion of the MOST system both in laboratory scale devices (ca. 5 cm²) using artificial sun, as well as in larger scale (ca. 900 cm² irradiated area) devices under natural sunlight, outdoor conditions. This is, to our knowledge, the first attempt to bring the MOST system out of the lab, demonstrating one possible avenue for future lab-to-site transfer. Importantly, this experiment verifies that DHA can be photochemically converted both under

simulated conditions and under natural outdoor conditions. The experiments are accompanied by a discussion seeking to identify where future optimization of the DHA/VHF couple should be focused.

Results and Discussion

An important feature of a MOST system is its ability to absorb photons efficiently, store the energy for extended periods of time, and recover the energy upon thermal back-conversion. These processes should ide-

ally proceed without degradation. Figure 1 shows the absorption spectra of DHA1 dissolved in toluene after exposure to an increasing amount of light, showing clear isosbestic points, and thus a clean conversion into VHF1. The robustness of the system may vary in different solvents, and the rate of the thermal back-conversion of VHF to DHA is known to be very solvent dependent; being faster in more polar solvents.[12] Three candidate solvents were selected for testing in order to cover a range of important solvent characteristics; namely, toluene (nonpolar), acetonitrile (polar aprotic), and ethanol (polar protic). The solubility of the molecules is also a critical parameter, since together with the energy density of the pure compounds it determines the maximal adiabatic temperature gradient one can expect to achieve upon heat release.^[6f] The solubility of DHA1 in the three studied solvents was measured according to a slightly modified literature procedure^[13] (see the Supporting Information, section I for details; results summarized in Table 1). Clearly, an increase in the solubility of pure **DHA1** correlates with decreasing polarity of the solvent, and it has a solubility of 0.2 m in toluene. A low-polarity solvent such as toluene, therefore, could combine a large energy density (high solubility of DHA) with a long, sustained release of the stored energy (slower back-conversion of VHF).



Figure 1. Absorption spectra of **DHA1** (in blue) and its photoisomer **VHF1** (in red); inset shows color change of the compound in toluene before and after irradiation with $\lambda = 365$ nm light.

Table 1. Summary of physical properties of DHA1 in different solvents.					
Solvent	Photoisomerization quantum yield ^[14]	Thermally induced back- conversion half-life at 25 °C [min] ^[12]	Solubility [mg mL ⁻¹]	Degradation [% loss per cycle]	
toluene acetonitrile ethanol	0.6 0.55 0.5	1474 218 202	55 36 5	0.01 0.18 0.21	

Cyclability test

The robustness of the **DHA1/VHF1** couple was probed by subjecting solutions of the photoswitch to as many as 70 complete photothermal cycles. The characteristic absorbance of **VHF1** at $\lambda = 460$ nm was used to follow both processes, revealing a remarkable stability, with less than 0.01% degradation per cycle in toluene. It also demonstrates that **DHA1** degrades faster in polar solvents with around 0.28% and 0.18% loss of molecules per cycle for ethanol and acetonitrile, respectively (Figure 2).

The ethanol and acetonitrile solutions were analyzed by LC-MS after the cycling tests. Many components could be seen, and two of these seem to be azulenes resulting from either elimination of HCN from **DHA1** or a substitution reaction (presumably S_N2') through nucleophilic attack by EtOH followed by an oxidation (Figure 3) in accordance with previous observations.^[15]

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Figure 2. Cyclability test based on absorbance at $\lambda = 460$ nm in toluene (blue, ca. 10^{-4} M), acetonitrile (purple, ca. 10^{-4} M) and ethanol (red, ca. 10^{-4} M). To reduce the back-conversion time, the experiments were performed at a constant temperature of 60 °C. Light at $\lambda = 365$ nm was repeatedly switched on and off. One cycle corresponds to one light-on period and one light-off period. A total of 70 cycles are shown. The baseline absorption increase for the ethanol sample is attributed to the formation of azulene. Parasitic signals (spikes) appear in acetonitrile and ethanol samples due to instrumental fluctuations.



Figure 3. Possible degradation products based on mass spectrometry analysis.

Catalytic conversion of VHF1 to DHA1 in flow reactor

With a view to triggering the energy release on demand, we recently found that the rate of the back-reaction from VHF1 to DHA1 can be significantly enhanced by Cu¹ ions.^[10e] Thus, for VHF1 we found a decrease in the half-life from 545 min at 25°C in dichloromethane to 18–22 min upon addition of 1 equivalent of $[Cu(CH_3CN)_4]BF_4$. Here, we demonstrate that the catalyzed reaction can be carried out under flow conditions by using a Cu¹ salt in a fixed-bed flow reactor. First, a DHA1 solution (ca. 1.7×10^{-3} M) in toluene was fully converted into VHF1 by using a solar simulator lamp for 1 h. A flow reactor was constructed from Teflon tubing and packed with solid $[Cu(CH_3CN)_4]PF_6$ crystals. A flow speed of 5 mLh⁻¹ was selected to back-convert VHF1 into its isomer. Up to 81.3% of VHF1 was converted into DHA1 (see Figure 4 and Experimental Section for details).

Indoor microfluidic chip device

For energy storage applications, the ability of the DHA1/VHF1 couple to store energy is of prime importance. Calculations show a Gibbs free energy difference between DHA1 and VHF1 of 27.7 kJ mol⁻¹ and an enthalpy difference (ΔH_{stor}) of 35.2 kJ mol⁻¹. ΔH_{stor} values were used in the estimation of the solar energy storage efficiency η_{MOST} [as determined by Equations (1)–(3); see below].^{(10b,16]} From the point of view of molecular stability, toluene, or a similar nonpolar solvent, seems to be the most attractive choice for device testing, because of its long VHF half-life, high quantum yield, high solubility, and minimal degradation. Thus, toluene was employed in the device testing experiments.

To evaluate the basic function of the DHA1/VHF1 MOST system under continuous-flow irradiation conditions, a micro-



Figure 4. Catalytic conversion in flow reactor: a, b) Scheme and photograph of the back-conversion setup. Boxes 1 and 2 contain in-line UV/Vis spectrometers. c) UV/Vis spectra before (red) and after (blue) exposure to $[Cu(CH_3CN)_4]PF_6$.



fluidic reactor was built to test the conversion ratio of **DHA1** into **VHF1**. Conversion was tested at two different concentrations in toluene by using simulated sunlight (1 sun, AM1.5). Solutions of **DHA1** at 1×10^{-3} M and 4×10^{-3} M were pumped through a 85 µm optical path-length microfluidic chip, and the conversion was determined by monitoring the UV/Vis absorption using two flow-cell spectrometers positioned before and after irradiation (Figure 5).



Figure 5. a) Microfluidic chip test setup. Boxes 1 and 2 contain in-line UV/Vis spectrometers and box 3 contains the microfluidic chip. Green line shows optical waveguides containing split and combined probe light. b) Picture of the used setup.

The capacity of a MOST system to store energy can be characterized by η_{MOST} . Intuitively, the theoretical maximum of the energy storage limit is a function of the absorption overlap with the solar spectrum, concentration, photoisomerization quantum yield (ϕ_{iso}), and energy difference between the two isomers (ΔH_{stor}). The theoretical capacity is expressed in Equation (1):

$$\eta_{\text{MOST}} = \frac{\int_{0}^{\lambda_{\text{Onset}}} \frac{E_{\text{AM1.5}}(\lambda) (1-T) \phi_{\text{los}} \Delta H_{\text{stor}}}{h \nu N_{\text{A}}} \frac{d\lambda}{d\lambda} \times 100 \%$$
(1)

where $E_{AM1.5}(\lambda)$ is the spectral irradiance (energy current density) in Js⁻¹m⁻²nm⁻¹; *T* is the transmittance of DHA in specified device; *h* is the Plank constant in J s; ν is the frequency of incoming light in s⁻¹; and N_A is Avogadro's constant. The photoisomerization quantum yield is defined as in Equation (2):

$$\phi_{\rm iso} = \frac{\text{number of isomerization events}}{\text{number of photons absorbed}}$$
(2)

Based on idealized conditions, that is, neglecting reflection losses, the calculated maximum energy storage efficiency is 0.61% for a saturated solution of **DHA1** in toluene. Thus the maximum efficiencies for the two solutions investigated in the device could be at most 0.13% at 4×10^{-3} M and 0.04% at 1×10^{-3} M (Figure 6a). The experimental energy storage efficiency is calculated using Equation (3):

$$\eta_{\text{MOST}} = \frac{\dot{n}_{\text{DHA}} \, \alpha_{\text{VHF}} \, \Delta H_{\text{stor}}}{A \, E_{\text{AM1.5}}} \tag{3}$$

where \dot{n}_{DHA} is the **DHA1** flow speed in mols⁻¹, α_{VHF} is the conversion ratio of **DHA1** into **VHF1** and *A* is the irradiated area in m².



Figure 6. a) Simulated maximum energy storage efficiency as a function of DHA concentration through a 85 μ m path-length device. The maximum concentration (0.21 M) corresponds to **DHA1**'s maximum solubility in toluene. Red and blue triangles correspond to the concentrations of **DHA1** used experimentally, 4×10^{-3} M and 1×10^{-1} M, respectively. b, c) Conversion percentage and evolution of the energy storage efficiency as a function of residence time in the microfluidic chip for these solutions (average taken from three separate measurements). In (c), red and blue triangles correspond to the calculated value from (a).

Experimentally, by controlling the flow speed, different fluid residence times were obtained ranging from 3 s to 43 s. The variation in residence time increased the conversion percentage of **DHA1** into **VHF1** from around 40% to a complete transformation for both used concentrations (Figure 6b). Interestingly, an irradiation of 10 seconds resulted in about 80% conversion in both cases. The maximum measured energy storage efficiency in the microfluidic chip was found to be 0.13% at 4×10^{-3} M and 0.034% at 1×10^{-3} M, at a residence time of 3 s, which are close to the theoretical limits (Figure 6c). To put



these numbers in context, solar energy conversion efficiencies have been reported under similar conditions for ruthenium fulvalene (Ru₂Fv) compounds (0.0071%) mainly limited by a poor quantum yield and norbornadienes (up to 1.1%),^[2,17] limited by quantum yield and onset of absorption.

a)

DHA1

 \mathbf{O}

An ideal MOST candidate molecule should show a quasiconstant efficiency at different concentrations and long residence times. Ideally, the metastable isomer should not absorb light in the solar spectrum. However, the energy storage efficiency decreases with longer residence times in the chip. This behavior is caused by an inner filter effect due to VHF absorbing light in the visible and UV range of the solar spectrum, thus parasitizing on the incoming solar energy necessary for the photoconversion of DHA1 into VHF1.

Outdoor parabolic concentrator

In order to demonstrate the function of the system under real outdoor conditions, a parabolic concentrator combined with a solar tracker was constructed for outdoor testing. This device also works as a proof-of-principle experiment toward future lab-to-site transfer. Sun light was focused on an apparatus consisting of jacketed glass tubing (Figure 7).^[6a] The inner tube serves as a channel for cooling water to avoid back-conversion caused by an increased temperature from the focusing light. The DHA1 solution was pumped through the 5.8 mL outer jacket to maximize the direct exposure to sunlight. Importantly, this heating of the water can be exploited as an auxiliary product in hybrid water heating/MOST devices, as we recently reported.^[17]

In this case, a saturated solution of DHA1 can give a theoretical limit storage efficiency of 0.65% in toluene. For this experiment, a 3.7×10^{-3} M toluene solution of **DHA1** was pumped through the system at variable residence times, ranging from 35 s to 417 s (ca. 600 to 50 mLh⁻¹), giving conversion percentages from 84% up to 100%. However, estimation of the efficiency by using Equation (3) and assuming that the solar flux was the standard AM1.5, the measured maximum energy storage efficiency was only 0.02%, which is far from the theoretical maximum of 0.45% (Figure 8). Since the microfluidic device gave a good match between calculated and measured efficiencies, we accredit the majority of the large discrepancy found in the scaled-up device to a much larger device volume and a longer path length consequently amplifying the inner filter effects from competition of absorption between DHA1 and VHF1, as well as a lower actual solar influx than the standard AM1.5 and minor misalignment of the concentrator. Future device and molecular designs should take the challenge with inner filter effects into account by having shorter path lengths, redshifting the strong VHF absorbance or lower spectroscopic overlap between the isomers.

Conclusions

In this study, we have evaluated the DHA1/VHF1 photothermoswitch for MOST applications. Initial cycling experiments showed that a nonpolar solvent (toluene) provides a larger sol-



Figure 7. Outdoor parabolic concentrator setup: a) Schematic of the device. b) Actual working design. Boxes 1 and 2 represent the flow spectra detectors, as in Figure 4. Box 3 is the two-layer glass tube and box 4 shows the parabolic concentrator. Cooling water serves to keep the MOST system at sufficiently low temperature.

ubility and less degradation for long-term use than polar solvents. Moreover, the [Cu(CH₃CN)₄]PF₆-catalyzed conversion of VHF into DHA was demonstrated in a flow reactor. We also tested the solar energy storage in a laboratory-scale device, as well as, for the first time, under realistic outdoor conditions. The controllable microfluidic chip shows a good energy storage efficiency, which is very close to the theoretical estimate. Concerning the outdoor parabolic device, a need was identified for better molecular candidates with lower spectral overlap between the two isomers. Based on theory, future improvement can be focused on the enhancement of the energy storage capacity of the DHA/VHF couple (ΔH_{stor}), which is only 0.14 MJ kg⁻¹ (calculated for neat compound),^[16] but, by structural modifications such as removing one of the two cyano groups $^{[10b]}$ or by fusing a benzene ring to the DHA, $^{[10f,g]}$ can be increased 2-4 times. However, these modifications have to date had undesired implications for the switching and stability



properties, and it remains a challenge to find the optimum molecule satisfying all of these requirements. Moreover, we note that, since the use of solvents decreases the possible obtainable heat gradient, future molecular design should focus on increasing solubility, or on developing solvent-free systems.^[6f,g] Potentially, it could be envisaged that a scale up of this system could be combined with solar water heating^[17a] or photon up-conversion^[6a] to increase the utilization of incoming photons. Finally, we envision the devices demonstrated herein to be applicable for other types of photoinduced chemical processes, such as photoinduced redox chemistry for green chemical production.[18]

Experimental Section

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a)

storage efficiency (%)

energy s

Conversion (%) 90

80

70

Gothenburg, Sweden.

b)

0.00

0.05

110

0.10

220

Simulated maximum

General: All solvents were HPLC grade. EtOH used was 99.5% in purity. Solutions of DHA1 were kept in the dark during preparation, storage, and handling until exposure to light in the testing devices (microchip or glass tube).

Determination of the solubility of DHA in PhMe, MeCN, and **EtOH**

General: A modified version of a previously reported procedure was adopted. $^{\scriptscriptstyle [13]}$ For internal standard, 1,4-dinitrobenzene (>99% purity) was chosen.^[19] The employed DHA1 (2-phenylazulene-1, 1(8aH)-dicarbonitrile) was recrystallized from boiling n-heptane (HPLC grade) to >99% purity according to ¹H NMR spectroscopy. All weightings were performed on a Sartorius Extend ED2245 d=0.1 mg analytical balance. All quantitative ¹H NMR (qH NMR) experiments were performed at 298 K on a Varian 400/54 spectrometer operating at 399.94 MHz for ¹H. The total time occupied on the NMR instrument (including overhead for sample changes, locking, shimming, etc.) for the entire range of measurements in this investigation was roughly 9 h. Chemical shifts are referenced to CHCl₃ (δ = 7.26 ppm).^[20]

Solubility determination by qH NMR spectroscopy: Saturated solutions of DHA1 in the respective solvents were prepared by thorough mixing for 24 h. Aliguots were taken, spiked with standard solution, and the resulting samples investigated by ¹H NMR spectroscopy under quantitative experimental conditions. Selected signals were carefully integrated to give molar ratios of the sample constituents, that is, the standard, DHA, and solvent. The molar ratios were then converted into solubility figures.

Cyclability test

A quartz cuvette with four transparent open-access sides was fixed on a magnetic stirrer, then covered by a water heating system, and inserted into a spectrophotometer (Agilent Cary 100 UV/Vis for toluene sample, Agilent Cary 4000 UV/Vis for ethanol and acetonitrile sample). Three solutions, with concentrations of around $0.6 \times$ 10^{-4} M, 0.5×10^{-4} M, and 0.6×10^{-4} M, were prepared in toluene, acetonitrile, and ethanol, respectively. Two of the three samples were degassed by nitrogen flushing. However, due to the low boiling point of ethanol, that corresponding sample was prepared by the freeze-pump-thaw technique (ca. 5 cycles) to eliminate dissolved oxygen. A $\lambda = 365$ nm collimated LED light source (Thorlabs) controlled by a custom-made LabVIEW program was used to irradiate the samples. The irradiation times were 60 s in toluene, 300 s in EtOH, and 200 s in MeCN, and the corresponding heating times were 6500 s, 1400 s, and 2300 s, respectively. The heating system was monitored at 60 $^\circ\text{C}$ during the whole experiment with a maximum fluctuation of +/-4 °C. The recorded signal was obtained from a $\lambda = 460$ nm probe light close to the maximum absorption peak of VHF in the three solvents. All photoisomerizations and thermally induced back-conversions were monitored over at least 80% of the observed absorbance.

Catalytic conversion of VHF1 to DHA1 in flow reactor

[Cu(CH₃CN)₄]PF₆ (10 mg; 97% purity, Aldrich) was loaded into a Teflon tube (1.27 mm inner diameter, 2.5 cm long; Cole-Parmer Instrument Company). Two pieces of cotton were used to prevent the copper salt from exiting the reactor. The spectrum before and after conversion was recorded by using two flow-cell spectrophotometers (Avantes, AvaSpec-2048L), placed before and after the catalytic reactor, with path lengths of around 0.03 and 0.09 cm for each channel individually.

Microfluidic chip

The microfluidic chip made from fused silica was acquired from Syrris Ltd. with a total volume of 62.5 µL, an effective exposure volume of 35.72 µL, irradiation surface of 5.18 cm², and a channel depth of 85 µm. The fluid speed was controlled by a pump from the same source. A solution of DHA1 with concentration of $4\times$ 10^{-3} M was prepared and stirred in the dark over 1 h. The path length of flow cells were determined to be around 0.06 and 0.10 cm for each channel individually.

Parabolic concentrator

Into a 500 mL volumetric flask, DHA (475.4 mg, 1.855 mmol), and toluene (ca. 100 mL) were added, after which the flask was swirled