Acylhydrazone-based supramolecular assemblies undergoing a converse sol-to-gel transition on trans $\rightarrow$ cis photoisomerization†

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Photoisomeric supramolecular assemblies have drawn enormous attention in recent years. Although it is a general rule that photoisomerization from a less to a more distorted isomer causes the destruction of assemblies, this photoisomerization process inducing a converse transition from irregular aggregates to regular assemblies is still a great challenge. Here, we report a converse sol-to-gel transition derived from the planar to nonplanar photoisomer conversion, which is in sharp contrast to the conventional light-induced gel collapse. A well-designed acylhydrazone-linked monomer is exploited as a photoisomer to realize the above-mentioned phase transition. In the monomer, imine is responsible for the planar to nonplanar photoisomer conversion, which is in sharp contrast to the conventional light-induced gel collapse. A well-designed acylhydrazone-linked monomer is exploited as a photoisomer to realize the above-mentioned phase transition. In the monomer, imine is responsible for the planar to nonplanar photoisomer conversion, which is in sharp contrast to the conventional light-induced gel collapse. A well-designed acylhydrazone-linked monomer is exploited as a photoisomer to realize the above-mentioned phase transition. In the monomer, imine is responsible for the planar to nonplanar photoisomer conversion, which is in sharp contrast to the conventional light-induced gel collapse.

Introduction

Photoisomerization is a process that reversibly converts one isomer into another with a clean and remote light source, which has been applied in a broad range of systems, including photoactuators, sustained drug release systems, information storage, photochromic materials, and photogated devices.1-9 Photoisomeric supramolecular assemblies, as an elegant combination of reversible photoisomers and dynamic noncovalent interactions, possess various intriguing properties, such as degradability, shape memory, and phase transition.10-14 Generally, photoisomerization from a relatively planar isomer to a distorted one gives rise to the disassembly of supramolecular systems, which is ascribed to the steric hindrance of the nonplanar isomers failing to orderly stack with each other (Fig. 1A). For instance, based on trans $\rightarrow$ cis photoisomerization, azobenzene-containing molecules induced a solid to isotropic liquid phase transition,15-18 and cyanostilbene derivatives caused not only a liquid crystal to an amorphous phase transition,19-20 but also a gel collapse (gel-to-sol transition).21 Nevertheless, although a few examples of transition from a less to a more ordered state upon photoirradiation have been reported in liquid crystal systems, the trans $\rightarrow$ cis photoisomerization-induced converse transition from irregular aggregates to regular assemblies is far less reported so far and still a great challenge (Fig. 1B). This is mainly because the bent-shaped (cis)-isomers loosely stack and consequently lead to weaker intermolecular interactions. However, this converse transition could enrich the categories of supramolecular photochemical phase transitions. Moreover, the

Fig. 1 (A) The conventional self-assembly behaviors of trans and cis monomers, and (B) the converse self-assembly and the transition on trans $\rightarrow$ cis photoisomerization.

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formation of light-induced regular assemblies (such as gelation) perfectly meets the requirements for in situ gelation in a sealed environment, such as in the microfluidics field. Thus, the development of ingenious methods to enable the formation of light-induced regular assemblies from trans → cis conversion is of pivotal importance for both a fundamental viewpoint and potential applications.

An effective strategy to realize the above photoisomerization-induced transition from irregular aggregates to regular assemblies is to seek photoisomers with unique structures. Acylhydrazones, 24–27 a novel class of photoisomers based on the imine structural unit (E–Z interconversion of the C==N bond), 28–30 have recently been regarded as promising candidates for photo-switches because of their facile synthesis, high stability, and fatigue resistance. More importantly, di-arylethers are abbreviated as ethoxy groups to reduce computational costs. The amide in acylhydrazones is an excellent fragment to generate a hydrogen bond, which is known as a strong driving force for constructing regular assemblies. 38–40 On the other hand, the aggregated nature of supramolecular assemblies usually causes incomplete photochemical conversion, 41–43 further leading to the coexistence of E- and Z-form isomers in the final state of supramolecular systems, which makes it possible to manipulate the stacking mode of the two isomers. Benefitting from all these essential advantages, we envision that, acylhydrazone-functionalized monomers could be exploited to realize the geometrical isomerism induced converse assembly transition on trans → cis conversion (Fig. 1B), in a supramolecular co-assembly manner by photo-regulating the balance between isomerization and aggregation.

Based on these considerations, we herein realize a converse sol-to-gel transition derived from the trans → cis photoisomerization of a low-molecular-weight acylhydrazone-linked monomer (Fig. 2), which is in sharp contrast to the previously reported light-induced gel collapse. 44,45 Specifically, a tadpole-like monomer 1 consisting of a pyrene head and hydrophobic tail linked by an acylhydrazone unit, is first designed and synthesized. It can undergo a reversible E–Z photoisomerization process in both the monomeric and assembled states. Upon irradiating the concentrated decane solution of 1-E, a sol-to-gel transition is realized. This phase transition is ascribed to the co-assembly of 1-E and partial photoisomerization product 1-Z, mainly driven by the emergence of intermolecular hydrogen bonds between the two adjacent monomers. On this basis, the light-induced sol-to-gel transition is successfully employed to establish the valves of microfluidic channels based on a reversible and straightforward gelation process.

Results and discussion

Photoisomerization of the acylhydrazone-linked monomer

We first investigated the photoisomerization behavior of 1 in the monomeric state. In dilute dichloroethane (DCE, c = 5.0 × 10⁻³ M), the UV-vis spectra of the pure (E)-isomer of 1 exhibited vibronically split absorption bands in the region of 325–425 nm (Fig. 3A and ESI Fig. S1†). Upon irradiation with 365 nm light for 50 s, the maximum absorption wavelength (λmax) of 1-E blue-shifted from around 374 nm to 346 nm with concomitant two isosbestic points at 315 and 351 nm, indicative of the photoisomerization from 1-E to the photostationary state (Z)-isomer of 1 (1-ZPSS). Irradiation of 1-ZPSS with a shorter wavelength light of 254 nm regenerated the initial (E)-isomer, which was confirmed by the complete restoration of the absorption bands (ESI Fig. S1†). With successive 365 nm and 254 nm light irradiation, the photoisomerization of 1 could be reversibly

![Fig. 2](image-url)  
Fig. 2 Schematic illustration of the photoisomerization of monomer 1, and a cartoon representation for the proposed self-assembly behaviors of 1-E and 1-Z, and its trans → cis photoisomerization induced supramolecular sol-to-gel transition.

![Fig. 3](image-url)  
Fig. 3 Photoisomerization studies of monomer 1. (A) UV-vis spectral change of 1-E (5.0 × 10⁻³ M in DCE) upon successive 365 nm (irradiation intensities: 0.98 mW cm⁻²) and 254 nm (irradiation intensities: 0.34 mW cm⁻²) light irradiation. Inset: multicycle experiments by monitoring the absorbance intensity changes at 373 nm. UV-vis spectral change of the control compound 2 (5.0 × 10⁻³ M in DCE) upon 365 nm light irradiation. (B) Partial 1H NMR spectra (CD₂Cl₂, 298 K, 2.00 mM) of 1-E: (i) before irradiation, (ii) conversion to 1-ZPSS after 365 nm light irradiation for 60 min, and (iii) recovery to 1-E upon 254 nm light irradiation for 30 min. (C) The calculated energy levels and corresponding frontier molecular orbital diagrams of 1-E and 1-Z via DFT computation. (D) The side view and top view of the optimized structures and related Gibbs-free energy of 1-E and 1-Z based on DFT calculations at the B3LYP/6-311G(d) level. The long alkyl chains in 1-E and 1-Z are abbreviated as ethoxy groups to reduce computational costs.
switched multiple times without noticeable degradation in performance (Fig. 3A and ESI Fig. S2†). Furthermore, based on the quantitative analysis of typical 1H NMR resonances of 1-E in C2D2Cl4, 1-E to 1-ZPPS photoisomerization was about 57% complete, while the back-isomerization was almost entirely complete within 30 min (>98% yield, Fig. 3B). The time-dependent absorbance curve conformed to the pseudo-first-order kinetics, providing an E-Z photoisomerization rate (kobs) of 0.058 s−1 (ESI Fig. S3†). kobs was related to the concentration of 1-E, which increased as the concentration increased. In addition to 254 nm light irradiation, heating could also trigger the back-isomerization (ESI Fig. S4†). In comparison, the reference compound 2 (varying acylhydrazine to ester bond) was unaffected by light irradiation, as evidenced by the complete lack of changes in the UV-vis and fluorescence signals (Fig. 3A, inset, and ESI Fig. S5†).

To get further insights into the relationship between acylhydrazine photoisomerization and molecular structures, quantum chemical calculations on both isomers of 1 were employed. Based on the comparative analysis of the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) of 1-E and 1-Z in the ground state (Fig. 3C), the calculated energy gap of 1-E (3.29 eV, λ = 376 nm) was relatively lower than that of 1-Z (3.56 eV, λ = 348 nm), which were consistent with the λmax values measured by the UV-vis spectra. The HOMO and LUMO electron densities were localized at the pyrene moieties for 1-E, reflective of the transition from π to π* orbitals. The similar phenomenon was observed for 1-Z. For the optimized structures of 1-E and 1-Z based on density functional theory (DFT) calculations at the B3LYP/6-311G(d) level, the total energy (E) values were calculated to be −1570.457880 and −1570.452184 a.u., respectively (Fig. 3D). It was found that 1-E was more stable with approximately 15.0 KJ mol−1 lower energy relative to 1-Z, because of the more twisted molecular structure of 1-Z.

Light-induced sol-to-gel transition

The assembly behaviors of monomer 1-E before photoirradiation were first studied. As is well known, hydrocarbon solvents such as decane are the common poor solvents for rod-coil-type monomers to form ordered supramolecular assemblies.47-49 When switching the solvent from DCE to decane, the UV-vis absorption spectra of 1-E showed an emergence of a shoulder band located between 400 and 475 nm (ESI Fig. S6†). Simultaneously, the intensity of the blue emission band at 412 nm corresponding to the monomeric pyrene units slightly decreased, along with the appearance of a new emission band at 520 nm (ESI Fig. S6†). The red-shifted structureless band was attributed to the excimer emission of the stacked pyrene units.50-52 Unfortunately, no noticeable aggregation of 1-E could be detected by the Tyndall effect experiment, and dynamic light scattering (DLS) and transmission electron microscopy (TEM) measurements (ESI Fig. S7†). Even on increasing the concentration up to 20 mM, no gels were visualized during the heating-cooling cycle, and remained unchanged in a dark place for several days (ESI Fig. S8†). Accordingly, no large-sized supramolecular assemblies were formed for 1-E in decane.

Surprisingly, when the concentrated decane solution of 1-E was exposed to 365 nm light, the viscosity gradually increased and fluidity was lost. Subsequently, a yellow transparent gel was finally formed within 60 min (Fig. 4A and ESI Fig. S9†). Such results are in sharp contrast to the previously reported light-induced gel collapse.21,44,45,53 To acquire the detailed gelation process, viscosity experiments were then performed. Upon irradiating the decane solution of 1-E (10.0 mM), the specific viscosity was almost unchanged for the first 1.5 min and then increased sharply (Fig. 4B), indicating the favorable transition from monomeric species to higher-molecular-weight supramolecular assemblies. By varying the concentration, the viscosity transition time and intensity were finely adjusted. The critical gelation concentration was measured to be 6 mM. DLS analysis for 1-E under light irradiation showed the size enlargement phenomenon, revealing the light-induced supramolecular assembly behaviors (ESI Fig. S10†). In sharp contrast, no apparent viscosity and DLS hydrodynamic diameter (Dh) changes occurred for the reference compound 2 (Fig. 4B and ESI Fig. S10†). Furthermore, the light-induced sol-to-gel transition of 1-E was observed by microscopic morphologies. Depending on the TEM images, short nanofibers were observed for 1-E after irradiation for 10 min (Fig. 4C and ESI Fig. S11†). After 30 min, longer fibers with several microns in length were formed. It is thus envisioned that these one-dimensional fibers are mutually entangled to create three-dimensional fibrous networks that thereby entrap a large number of solvent molecules, which lays the basis for the formation of supramolecular gels.

The properties of the light-induced gel were further proved by means of rheological experiments. The storage modulus (G′) value was remarkably higher than the loss modulus (G″) value at all tested frequencies (Fig. 4D), which verified the formation of

![Fig. 4](light-induced_sol-to-gel_transition_of_1-E.png)
true supramolecular gels. Additionally, the resulting organogels were stable enough once they were formed. The conclusion was confirmed using the oscillation time curves, indicating that the values of $G'$ and $G''$ remained constant over a long time ($\sim 30$ min, Fig. 4E). The stacking pattern of the light-induced gel was identified by small-angle X-ray diffraction (XRD) measurements. Specifically, no regular stacking formed for 1-EE before irradiation, as evidenced by the lack of reflection signals in the XRD pattern (Fig. 4F). Upon irradiating 1-EE with 365 nm light, the diffractogram showed three reflections with d-spacing values in the ratio of $1/\alpha : 1/\beta : 1/\gamma$, assigned to the (100), (110), and (200) signals of the hexagonal columnar (CoH$_6$) stacking. The lattice parameter ($\alpha = 2d_{100}/\gamma = 3.78$ nm) calculated from the diffraction maxima indicated that the diffraction units are comparable to the stacking model diameter of light-irradiated 1-EE (3.89 nm, Fig. 4F, inset, and ESI Fig. S12†). Additionally, thermal stimulus ($\sim 45$ °C) could also trigger the liquefaction process, and the sol remained even on cooling to room temperature.

For the purpose of investigating the stability of light-induced organogels from the thermodynamic standpoint, we turned to elucidate the assembly mechanism of the supramolecular assemblies. The organogels were diluted to an appropriate concentration for spectral measurements. Temperature-dependent UV-vis spectra were then recorded. In detail, upon gradually decreasing the temperature at a rate of 60 K h$^{-1}$, the absorption spectra of the diluted gel of light-irradiated 1-EE showed two isosbestic points at 300 and 399 nm (Fig. 5A, inset), indicative of the transition between monomeric and assembled states. A non-sigmoidal cooling curve was obtained by monitoring the fraction of aggregated species ($\alpha_{agg}$) against temperature (Fig. 5A and ESI Fig. S14†), suggesting that the supramolecular assembly process followed a cooperative nucleation–elongation mechanism. The Meijer–Schenning–van-der-Schoot mathematical model was then employed for the non-linear fitting of the resulting curves. A $T_c$ (critical elongation temperature) value of 310.4 K and a $K_a$ (dimensionless equilibrium constant of the activation step at $T_c$) value of $8.6 \times 10^{-5}$ were obtained. Moreover, on the basis of the modified Van’t Hoff plot, $\Delta G^\circ$ (Gibbs free energy) was calculated to be $-26.2$ kJ mol$^{-1}$ at 298 K (Fig. 5B). Such results manifest that organogels derived from light-induced supramolecular assemblies possess considerable stability.

**Insights into the sol-to-gel transition mechanism**

For a deeper understanding of the origin of the gelation behaviors triggered by photoirradiation, UV-vis spectra of 1-EE (5.0 $\times$ $10^{-5}$ M) in decane upon exposure to 365 nm light were first recorded. In detail, the original absorbance signal decreased in intensity, while a blue-shifted band centered at 345 nm gradually turned into the main band, with two isosbestic points at 322 and 353 nm (ESI Fig. S16†). The results implied that the trans $\leftrightarrow$ cis photoisomerization occurred not only in the monomeric state (DCE solution), but also in the assembled state. Due to the steric hindrance effect of stacked pyrene units, $k_{obs}$ in decane was slower than that in DCE (0.034 s$^{-1}$ versus 0.058 s$^{-1}$, ESI Fig. S17†). It is noteworthy that the trans $\leftrightarrow$ cis conversion of 1-EE at a relatively higher concentration cannot reach the conversion level at a lower concentration, even on prolonging the irradiation time (ESI Fig. S16†). Approximately 40% relative conversion was achieved at 0.2 mM compared to the lower concentration condition (Fig. 6A). Thus,

**Fig. 5 Supramolecular assembly mechanism studies.** (A) $\alpha_{agg}$ as a function of temperature for light-irradiated 1-EE in decane (5.0 $\times$ $10^{-5}$ M, $\lambda = 367$ nm). The solid lines denote the respective fitting curves. Inset: temperature-dependent UV-vis spectra. The arrows represent the signal variations upon decreasing the temperature. (B) Van’t Hoff plot for light irradiated 1-EE in decane. The red line denotes the linear fitting curve.

**Fig. 6 Insights into the light-induced sol-to-gel transition mechanism of 1-EE.** (A) The relative $E \leftrightarrow Z$ conversion of 1 (black dots: 5.0 $\times$ $10^{-5}$ M, red dots: 2.0 $\times$ $10^{-4}$ M) in decane upon 365 nm light irradiation, by monitoring the absorbance changes at 379 nm. (B) DLS hydrodynamic diameter variations upon gradual addition of 1-Z$_{PSS}$ (2.0 $\times$ $10^{-4}$ M) into the freshly prepared decane solution of 1-EE (2.0 $\times$ $10^{-5}$ M) and pure decane solvent, respectively. (C) SEM image of 10 mol% 1-Z$_{PSS}$ in 1-EE (2.0 $\times$ $10^{-4}$ M in decane). Insets display the enlarged P- and M-type helical fibers. (D) Optimized structures with color-mapped IGM iso-surface graphs of octamers of 1-EE (i) before, and (ii) after 365 nm light irradiation. The long side chains are simplified to ethoxy groups. The green color isosurfaces represent weak non-covalent interactions (such as van der Waals and $\pi-\pi$ stacking interactions), while blue color isosurfaces in (ii) denote stronger attraction (such as hydrogen bonds). The color scale at the bottom represents the relative strength of non-covalent interactions.
we rationalized that the supramolecular gel formation is ascribed to the partial photoisomerization of 1-E, that is, the participation of 1-Z.

We executed the titration experiments to further investigate the effect of light-irradiated product 1-Z_{PSS} on the formation of supramolecular co-assemblies. Upon progressively adding 1-Z_{PSS} into the decane solution of 1-E (2.0 × 10^{-4} M), the D_{h} value of assemblies determined by the DLS measurements gradually increased [Fig. 6B and ESI Fig. S18]. Only 10 mol% of 1-Z_{PSS} increased the D_{h} value from around 0.6 nm to 825 nm. As a comparison, no apparent enhancement of the D_{h} value was visualized when 1-Z_{PSS} was introduced into the pure decane solvent [Fig. 6B and ESI Fig. S18]. It thus indicated that the large-sized aggregates were formed by the co-assembly of 1-E with its light-irradiated product 1-Z_{PSS}. The conclusion was further verified by the morphology measurements. Intertangled long fibers were observed in the TEM images when adding 1-Z_{PSS} into 1-E (ESI Fig. S19), similar to that of 1-E exposed to 365 nm light for 30 min (Fig. 4C). Interestingly, the enlarged and scanning electron microscopy (SEM) images of the co-assemblies 1-Z/1-E showed that the fibers appeared as distinct right-handed (P) and left-handed (M) helical structures (Fig. 6C and ESI Fig. S19–S21). Therefore, it is envisioned that the structurally distorted 1-Z displays a considerable capacity to regulate the dynamic stacking structures of the co-assemblies 1-Z/1-E to form well-defined supramolecular helical aggregates.

The detailed stacking mode for the light-induced supramolecular assemblies was then sought to elucidate. Depending on the 1H measurements, the NH protons of acylhydrazone displayed upfield and downfield shifts upon 365 nm light irradiation (ESI Fig. S22†). The significant upfield shift from NH to NH’ was assigned to the conversion from 1-E to 1-Z_{PSS}. The downfield shift of the unisomerized NH proton revealed the presence of hydrogen bond interactions after photoirradiation. The hydrogen bonds were then verified through Fourier transform infrared (FTIR) spectra experiments. The N–H stretching vibration signal shifted from 3391 and 3195 cm^{-1} to a lower frequency (3172 cm^{-1}) upon irradiating 1-E with 365 nm light (ESI Fig. S23†), while the C=O band shifted from 1643 cm^{-1} to 1641 cm^{-1}, indicating the generation of intermolecular hydrogen bonds between neighboring acylhydrazone C=O and N–H bonds. Moreover, proton competition experiments were performed to evaluate the significance of hydrogen bonds for the light-induced supramolecular gel formation. Upon adding trace amounts of DMF into the gel system, the gel-to-sol transition was observed (ESI Fig. S24†), manifesting that the intermolecular hydrogen bonds were disturbed by the protic solvent of DMF, and therefore no fibrous structures formed under this condition.

To visually observe the non-covalent interaction variations upon partial photoisomerization from 1-E to 1-Z, the independent gradient model (IGM) methods of theoretical calculations were applied. The optimized structures with color-mapped IGM isosurfaces of the octamers of 1-E before and after photoirradiation are shown in Fig. 6D. In the IGM analysis, large-scope green isosurface regions existed between the pyrene moieties of the interdigitated self-assembly of 1-E, which was attributed to the π–π interactions (Fig. 6D(i)). After photoirradiation, the stacking structure transformed from the interdigitated way into the head-to-head manner. The emerged hydrogen bonds (Fig. 6D(ii), blue isosurface in the red frame) as well as π–π stacking and van der Waals interactions between 1-E and 1-Z synergistically stabilized the light-induced supramolecular assemblies. The average binding energies between the adjacent monomers increased from −7.69 kcal mol^{-1} of 1-E to −20.79 kcal mol^{-1} of light-irradiated 1-E.

Overall, these experimental and theoretical results unambiguously evidence that supramolecular gel formation is ascribed to the partial photoisomerization of 1-E causing the changes in stacking mode of the co-assembly 1-Z/1-E, mainly driven by the intermolecular hydrogen bonds.

**In situ sol-to-gel transition generated valves**

Microfluidic technologies have been recently regarded as an advanced tool to investigate non-equilibrium assemblies and controlled reactive processes in self-assembly research. Introducing stimuli-responsive materials as a valve would realize the non-contact microfluidic flow control, probably producing simplified and versatile microfluidic devices. Nevertheless, most reported valves were based on temperature- and electro-responsive systems, which are not easy to achieve flow control in situ and are comparatively complicated. Considering that the organogels derived from the light-induced sol-to-gel transition of 1-E can be finely triggered by remote and contactless light, they may be potential candidates for developing in situ generated valves in the microfluidics field.

On this basis, a three-layer microfluidic device was designed and fabricated, in which two circular chambers in the Y-shaped microchannel of the fluidic layer were used to fill the light-induced gel (Fig. 7A and B). Before employing 1 as the light-responsive valve, the decane fluid could pass through both directions of the microchannel (Fig. 7C(i)). The left side of the sealed microchannel was first filled with the pre-prepared high viscosity sol of 1 by injection, and the gel-based valve was then created by irradiating the sol with 365 nm light in situ. Thus, the left microchannel was blocked and the fluid could only pass through the right side (Fig. 7C(ii)). The fluid could be well maintained for several hours (even upon increasing the flow rate to around 20 µL min^{-1}), suggesting that the valve possesses enough stability. When the gel-based valve was irradiated with 254 nm light, the gel-to-sol transition occurred and then the sealability weakened, resulting in the recovery of bidirectional flow (Fig. 7C(iii)). It requires only a small amount of the gel-to-sol transition to restore the flow.

Besides, it is worth noting that the kind of liquid can be varied from low polarity decane to high polarity water. As shown in Fig. 7D and ESI Fig. S25, a similar flow control was achieved in the water-based environment. More importantly, the gel-based valves featured sufficient stability and sustainability in aqueous solution (even in saline). Specifically, 1-E could also undergo the light-induced sol-to-gel transition when mixed with saline solution (Fig. 7E). The gel did not swell or shrink after being immersed or sandwiched in saline for two weeks (Fig. 7F).
and ESI Fig. S26†). G′ and G″ were determined to be 612.2 Pa and 140.5 Pa, respectively (Fig. 7G), which are comparable to the values of the freshly prepared gel (645.3 Pa and 151.3 Pa, Fig. 4D). The helical fibers were also maintained (Fig. 7G, inset). Thus, the light-induced gel-based valves provide efficient and flexible flow control in sealed microfluidic chips.

Conclusions

In summary, we have successfully established a converse light-induced sol-to-gel transition system derived from the trans → cis conversion of acylhydrazone-based supramolecular assemblies. The acylhydrazone-linked monomer 1 underwent a reversible photosomerization process in both the monomeric and assembled states. A stable organogel was obtained upon irradiating the decane solution of 1-E with 365 nm light. This sol-to-gel transition is ascribed to the co-assembly of 1-E and light-irradiated product 1-ZPSS, mainly driven by the emergence of intermolecular hydrogen bonds between the two adjacent monomers in a head-to-head arrangement manner. We anticipate that, like acylhydrazone derivatives, other suitable stimuli-responsive molecules may manipulate the distinctive phase transition. Furthermore, the reversible sol-to-gel transition has been appropriately applied in gel-based valves formed in situ in microfluidic devices. The current work not only highlights the fundamental understanding of photosomerization in supramolecular photochemistry, but opens up a new avenue towards remote-controlled miniature materials.

Data availability

Detailed synthetic procedures, analytical data, and computational methods are provided in the ESI.

Author contributions

Z. G. and W. T. conceived the idea for this article. F. Y. and L. S. performed the experiments, analyzed the data, and produced the artwork under the direction of Z. G. Y. H. contributed to the theoretical calculations. S. Q. and J. Z. helped in the SEM measurements. F. W. and S. W. discussed the results and commented on the manuscript. The article was revised by Z. G. and W. T.

Conflicts of interest

There are no conflicts to declare.

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