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*Published in:*
Chemistry of Materials

*DOI:*
10.1021/acs.chemmater.6b03460

*Published: 22/11/2016*

*Document Version*
Publisher's PDF, also known as Version of record

*Please cite the original version:*
Efficient Light-Induced Phase Transitions in Halogen-Bonded Liquid Crystals

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ABSTRACT: Here, we present a new family of light-responsive, fluorinated supramolecular liquid crystals (LCs) showing efficient and reversible light-induced LC-to-isotropic phase transitions. Our materials design is based on fluorinated azobenzenes, where the fluorination serves to strengthen the noncovalent interaction with bond-accepting stilbazole molecules, and increase the lifetime of the cis-form of the azobenzene units. The halogen-bonded LCs were characterized by means of X-ray diffraction, hot-stage polarized optical microscopy, and differential scanning calorimetry. Simultaneous analysis of light-induced changes in birefringence, absorption, and optical scattering allowed us to estimate that <4% of the mesogenic units in the cis-form suffices to trigger the full LC-to-isotropic phase transition. We also report a light-induced and reversible crystal-to-isotropic phase transition, which has not been previously observed in supramolecular complexes. In addition to fundamental understanding of light-responsive supramolecular complexes, we foresee this study to be important in the development of bistable photonic devices and supramolecular actuators.

INTRODUCTION

Creating order and complexity through self-assembly of simple molecular building blocks is an important area of contemporary science. Yet, it is the external control over the molecular order that brings about technological disruptions, as well exemplified by the success of liquid crystals in display industry. Liquid crystals (LCs) constitute perhaps the best platform for exploring the interplay between self-assembly, molecular alignment, and stimuli-responsiveness, as they combine order and mobility in a unique way. As a remote, highly controllable energy source, light is a particularly attractive stimulus to control the LC phases. Through incorporation of light-responsive units into LC systems, several schemes for photomodulation of LC alignment have been devised, the main application potential lying in tunable photonics, functional surfaces, and light-to-mechanical energy conversion.

Azobenzene unit is a particularly efficient molecular switch for controlling the molecular alignment in LC systems, because of its large and reversible shape change upon photoisomerization and the incompatibility of the bent cis-azobenzene configuration with calamitic mesophases. The focus of azobenzene-based LC research can be simplistically divided into two main directions. On one hand, there is a thrust to develop tunable materials whose properties (e.g., surface topography) can be rapidly and reversibly changed between the undisturbed and photomodulated states. On the other hand, several photonic applications would benefit from bistable photomodulation, i.e., temporally stable switching between two states, using distinct wavelengths that drive the reverse photoisomerization reactions. Various structures for bistable azobenzene photoswitches have been developed, and, among them, ortho-fluorinated azobenzenes appear particularly promising. Fluorination is also an effective strategy to tailor the anisotropy and viscoelastic properties of LCs and boost intermolecular interactions for the self-assembly of supramolecular LCs. Combining these design principles, here, we target a new family of azobenzene-based, fluorinated supramolecular LCs with predictable phase behavior and efficient light-control over the molecular alignment.

In supramolecular LCs, specific noncovalent interactions drive the formation of supramolecular mesogens, even when starting from non-liquid-crystalline building blocks. They are pertinent from the viewpoint of increasing the modularity and tunability of conventional LC systems and obtaining...
functionalities not met in single-component or doped LCs. The majority of works in the field have concentrated on the design and structural characterization of supramolecular LCs. Their photochemical control, on the other hand, has not received much attention, whereas detailed studies have been conducted in single-component and doped LC systems. Understanding the dynamics of the light-induced processes, as well as the conditions under which photoinduced phase transitions occur in supramolecular systems, are not only fundamentally important, but also provide important design principles for supramolecular functional materials. Targeting photoresponsive, fluorinated supramolecular LCs, halogen bonding is the noncovalent interaction of choice.

Halogen bonding is an attractive interaction between an electrophilic region associated with a halogen atom in a molecular entity and a nucleophilic site. The benefits of halogen bonding, such as the high directionality and tunability, have recently been recognized in the design of functional materials. In particular, halogen-bonded, azobenzene-containing co-crystals and (amorphous) supramolecular side-chain polymers have shown extremely interesting photoresponsive properties, even superior to their hydrogen-bonded counterparts. Since the first demonstration in 2004, several halogen-bonded LCs have been reported. Photosensitive halogen-bonded LC complexes were first reported by some of us in 2012, and, in 2014, Yu et al. demonstrated photoinduced phase transitions in complexes between azopyridines and molecular halogens. However, neither of these studies detailed the photochemistry of the halogen-bonded LCs, which is crucial in terms of understanding their potential use in, e.g., supramolecular photonic devices and photoactuators.

Herein, we bridge this gap and report a detailed study on a small library of neutral halogen-bonded LCs, whose LC-to-isotropic phase transitions can be efficiently controlled with light. We show that <4% of the mesogenic units in the bent cis-configuration suffice to induce a complete order-to-disorder transition of the materials. We also report a light-induced and reversible crystal-to-isotropic phase transition, which has not been previously observed in supramolecular complexes.

RESULTS AND DISCUSSION
The materials used in this work are shown in Figure 1. We assembled 15 dimeric co-crystals 1-m-2-n in which the photoactive alkoxyazobenzene molecules 1-m contain an iodo-tetrafluoroarene ring acting as halogen-bond donor moiety, while 4,4'-alkoxystilbazoles 2-n were used as halogen-bond acceptors. Details about the synthesis, characterization, and sample preparation of the 1-m molecules and of their complexes are given in the Experimental Section. Recently, a complex of 2-1 with a halogen-bond-donating azobenzene substituted with a dimethylamino group has been reported to yield interesting photoresponsive behavior. Herein, we replaced the dimethylamino group of the azobenzene with an alkoxy chain because (i) long alkoxy chains should increase the mesophase stability, compared to the dimethylamino group, and (ii) by decreasing the electron-donating ability of the substituent, the lifetime of the cis-form of the azobenzene should significantly increase.

The photochemical properties of 1-m were first studied in 10^{-5} M DMF solution, and were observed to be identical for all three azobenzenes. Efficient trans-cis isomerization is attested by a decrease of the n-π* band at 363 nm and an increase of the n-π* band at 436 nm (Figure 2). The photostationary states were deduced by using the Fischer’s method, and they comprised 88%, 84%, and 24% of cis-azobenzenes upon illumination with 365, 395, and 457 nm, respectively. Note that the absorption spectrum of 2-n partly overlaps with that of 1-m (Figure 2), and upon irradiation at 365 nm, the stilbazoles also undergo photoisomerization. Therefore, 395 nm was the wavelength of choice for this work, yielding efficient isomerization of the azobenzenes while leaving the stilbazoles unchanged. At 20 °C in DMF, the cis-isomers of the 1-m molecules are highly stable, as illustrated in the inset of Figure 2. The thermal relaxation behavior is exponential, and the characteristic lifetime of the cis-isomer is ca. 12 days for all three molecules, confirming the design assumption (ii) of the previous paragraph. On time scales significantly shorter than the characteristic lifetime, the molecules can be considered to be bistable, which can be attributed to fluorination of the halogen-bond-donating phenyl ring.

The 15 halogen-bonded co-crystals 1-m-2-n were obtained by slow room-temperature evaporation of chloroform solutions containing the starting materials 1-m and 2-n in 1:1 molar ratios, which take into account the pairing of the respective strongest halogen-bond donor (I) and acceptor (pyridyl N) sites. 

Figure 1. Studied halogen-bonded photosensitive LCs 1-m-2-n are formed by self-assembly of the halogen-bond donors 1-m and the acceptor stilbazoles 2-n.

Figure 2. Normalized absorption spectra of the 1-m and 2-n molecules, represented by 1-10 and 2-2, respectively. The figure shows also the photostationary spectra under illumination wavelengths of 365, 395, and 457 nm, and the calculated spectrum of the cis-isomer for the 1-10 molecule. Inset shows the thermal cis-to-trans isomerization of the molecules at 20 °C indicating a cis-lifetime of ca. 12 days.
supramolecular organization of the starting materials in the co-crystals (see Figure 3, as well as Figures S1–S7 in the Supporting Information).

1-m and 2-n are assembled into supramolecular dimers by short and highly directional I--N(pyr) halogen bonds (see Figure 3). The respective I--N distances and C--I--N angles are given as follows: 2.771(2) Å and 178.6° in 1-8-2-2, 2.792(1) Å and 174.3° in 1-10-2-1, and 2.817(2) Å and 174.5° in 1-12-2-1. The observed halogen-bond distances roughly correspond to a 20% reduction of the sum of van der Waals radii of I and N.43

The strength of such a halogen bond was calculated to be ~6.5 kcal/mol in the reported complexes, which is very consistent with previous calculations.34 The high directionality of the halogen bond determines that calamitic structures are obtained, with aromatic rings of the two modules nearly coplanar in 1-10-2-1 and 1-12-2-1, and orthogonal in 1-8-2-2. The complexes 1-10-2-1 and 1-12-2-1 also exhibit arene–perfluoroarene quadrupolar interactions between neighboring 1-10 and 1-12 molecules in different planes. Because of these quadrupolar interactions, CH–O interactions, and CH–N interactions, the halogen-bonded dimers pack head-to-tail in an antiparallel fashion. Residual CH–F interactions further stabilize the overall crystal packing. Finally, powder X-ray diffraction (PXRD) analysis (Figures S8–S10 in the Supporting Information) confirmed that the crystalline bulk samples have the same crystal structures, as determined on the analyzed single crystals.

The LC phase behaviors of the complexes were analyzed by hot-stage polarized optical microscopy (POM) and differential scanning calorimetry (DSC). All complexes exhibited enantiotropic LC phases, with the exception of 1-12-2-12, which was monotropic. Nematic phases (Figure 4, left) were common to all complexes, while smectic A (SmA) phases (Figure 4, right) were observed only for complexes with the longest alkyl chains at both ends. We emphasize here that none of the starting compounds is liquid-crystalline as pure material, which implies that the halogen bond between the azobenzene and stilbazole units survives in the mesophase and drives the liquid crystallinity of the supramolecular structures.

The mesomorphism of the halogen-bonded liquid crystals described above is similar to that of analogous stilbazole complexes reported earlier,38 showing nematic phases at short chain lengths and SmA phases with longer chains. Apart from the complexes with the shortest chain-length stilbazole, the transition temperatures reveal a destabilization of the nematic phases with increasing n (Figure 5), which is a common behavior for calamitic mesogens with two terminal chains. A similar picture emerges when considering the variation of the alkoxy chain length on the azobenzene derivatives. Interestingly, the nematic phase extends over a broad temperature range, even exceeding 30 °C for many of the complexes. By preparing mixtures of several complexes, we were able to further suppress the onset temperatures for the LC phases, compared to single supramolecular entities. For instance, a mixture of 1-12 with 2-2 and 2-12 (in molar ratio of 1.0:0.5:0.5) exhibited a crystal-to-nematic transition at 79.5 °C, which is more than 10 °C lower than that of the pure 1-12-2-2.

Because of the presence of the azobenzene molecules, the complexes are photoresponsive and undergo a fast and reversible isothermal LC-to-isotropic phase transition upon irradiation with ultraviolet (UV) light (395 nm, 40 mW/cm²). This is illustrated for 1-10-2-8 at 110 °C in the POM.
micrographs shown in Figure 6, obtained using a planar LC cell with a thickness of 2 μm. We note that (i) the complex 1-10-2-8 was used as a model system, and (ii) similar behavior is observed also for other complexes and over the entire LC temperature range. Prior to UV irradiation, the sample exhibited a uniformly colored bright image when viewed between crossed polarizers with the director axis set to ±45°, with respect to the polarizer/analyzer (Figure 6a), whereas a black image was obtained when the axes coincided. This indicated homogeneous alignment of the LC complexes. As an example, the order parameter of molecular alignment is 0.44 for the nematic complex 1-10-2-8 at 105 °C, as determined from polarized absorption spectra. Under illumination, the birefringence due to the anisotropic molecular alignment disappeared within <3 s (Figure 6b), indicating the efficient photoinduced phase transition into an isotropic phase. Further illumination, continued in total for 35 s, did not induce further changes into the micrographs. Upon ceasing the illumination, the LC phase reappeared as small domains after a significant delay (Figure 6c), and, with time, the domains recombined and oriented along the rubbing direction (Figure 6d). The small isotropic droplets visible in the bottom right micrograph disappeared upon longer recovery times. We attribute the small LC domains (Figure 6c) as well as the isotropic droplets (Figure 6d) to phase separation between trans-rich liquid-crystalline and cis-rich isotropic domains that recombine once the cis-azobenzene fraction is sufficiently low.

From the POM images, we can deduce that the complexes exhibit a sequence of different photoinduced processes, and that the LC-to-isotropic phase transition happens relatively fast. In order to further quantify the photochemical processes, we simultaneously monitored the changes in absorbance, birefringence, and optical scattering at different stages of the illumination cycle, as shown in Figure 6e. Such measurements allowed us both to monitor the dynamics of the photoinduced LC-to-isotropic phase transition and to correlate the photoinduced phase separation with the phase transition process. Furthermore, they provided us with an estimate on how much cis-azobenzene is needed to induce phase transitions. To the best of our knowledge, such an effort has not been previously undertaken in supramolecular LCs.

The high birefringence prior to irradiation indicates that the molecules are initially well-oriented, and the decrease in birefringence once the UV illumination is started indicates rapid disappearance of the LC phase, in response to light irradiation. The photoinduced phase transition is accompanied by a rapid increase in both the attenuation at 400 nm (coinciding with the absorption band of the azobenzene), and attenuation due to optical scattering monitored at 700 nm, where neither of the compounds absorbs. The peak in scattering (the red curve in Figure 6e) indicates that photoinduced phase separation to LC and isotropic domains also occurs upon LC-to-isotropic phase transition. The abrupt increase in the attenuation at 400 nm once the irradiation is started (the blue curve in Figure 6e) arises from the combination of increased scattering due to phase separation, and changes in the UV-vis absorption spectrum due to a red-shift of the π-π* absorbance band of the azobenzene, attributed to disrupted molecular packing under UV illumination. The scattering peak disappears within a few seconds, whereas the absorbance at 400 nm continues to decrease steadily due to trans-cis isomerization for the entire illumination duration, or until the photostationary state is reached. Once the irradiation is ceased (35 s), the absorbance starts to immediately recover as the azobenzene molecules relax back to the thermally stable trans-form. The thermal relaxation is exponential and has a half-life of ca. 51 s at 110 °C (while the half-life at room temperature in a dilute DMF solution is ~12 days). However, there is a significant delay of ~130 s before the birefringence starts to recover. The onset of birefringence recovery is accompanied by the phase separation process shown in Figure 6c, which, based on the scattering peak, is now significantly slower than upon starting the illumination as it is driven thermally, not by light.
Based on our observations, there is a clear threshold in the cis-azobenzene concentration that determines the onset of the photoinduced phase transition at a given temperature (Figure S11 in the Supporting Information). By determining the spectra of the trans- and cis-isomers at the isotropic state (a detailed description is given in the Supporting Information (Figures S12 and S13)), the absorbance measurements allowed us to estimate that the onset cis-fraction of azobenzene molecules for the isotropic-to-LC phase transition is 7.6% ± 0.4%. This corresponds to the cis-fraction needed for the full LC-to-isotropic transition. Therefore, we can conclude that, similar to azobenzene-doped LCs,52,53 the bent shape of the cis-isomer effectively disrupts molecular order also in supramolecular LCs. Considering the 1:1 (azobenzene:stilbazole) nature of the complexes, and the fact that the stilbazole units are not isomerized under the irradiation conditions used, the overall number of cis-isomers needed to achieve the complete LC-to-isotropic phase transition is <4%.

The richness of the photoinduced processes in the complexes under investigation is further highlighted by the fact that, in addition to LC-to-isotropic phase transitions, they also undergo reversible crystal-to-isotropic transitions under irradiation with UV light. Photoresponsive crystalline materials have recently attracted much interest from the viewpoint of energy conversion, actuation, reversible adhesion, and controllable gas adsorption.59-63 Photoinduced crystal-to-isotropic transitions have been studied in azobenzene-based tetragonal compounds, ionic crystals, and cyclophanes.54-56 However, no such reversible isopomerization has been previously reported in supramolecular complexes. Figure 7a shows a POM image of the complex 1-10-2-8 at 85 °C, i.e., 10 °C below the melting point and 30 °C below the clearing point of the complex. Upon illumination (395 nm, 40 mW/cm²), a clear and reversible Cr−I transition occurred within 30 s (Figure 7b). Upon ceasing the irradiation, recrystallization occurred in ca. 3 min, through a partial nematic LC phase (Figure 7c), and the end result was a homogeneous crystalline phase (Figure 7d). The significant delay and the partial nematic phase indicate that the effect is due to photoisomerization rather than a photothermal effect.

CONCLUSIONS

We have shown that halogen-bond-driven supramolecular liquid crystals (LCs) between alkoxy-substituted azobenzene and stilbazole molecules exhibit a rich variety of photoinduced processes, including photoinduced LC-to-isotropic and crystal-to-isotropic phase transitions. Simultaneous analysis of light-induced changes in birefringence, absorption, and optical scattering allowed us to conclude that <4% of the mesogenic units in the cis-form suffices to trigger the full LC-to-isotropic phase transition. To the best of our knowledge, this is the first quantitative analysis of the phase transition process in supramolecular liquid crystals. In our materials design, fluorination plays an important role as it both increases the strong and directional halogen bonding to happen. Our future goals are to (i) suppress the temperature at which the complexes exhibit liquid crystallinity, and (ii) devise cholesteric, phototunable LCs driven by halogen bonding. Overall, our results demonstrate the versatility of halogen-bonded complexes as functional liquid-crystalline assemblies and pinpoint their potential toward tunable photonic devices and, eventually, supramolecular actuators.

EXPERIMENTAL SECTION

The starting materials were purchased from Sigma–Aldrich. Commercial HPLC-grade solvents were used without further purification, except for acetonitrile, which was used as a solvent for the synthesis of azobenzenes, which was dried over CaH₂ and distilled prior to use. 1H, 13C, and 19F NMR spectra were recorded at room temperature on a Bruker AV400 or AV500 spectrometer, using CDCl₃ as the solvent. 1H NMR and 13C NMR spectroscopy chemical shifts were referenced to tetramethylsilane (TMS) using residual proton or carbon impurities of the deuterated solvents as standard references, while 19F NMR spectroscopy chemical shifts were referenced to an internal CFCl₃ standard. The attenuated total reflection Fourier transform infrared (ATR-FTIR) spectra were obtained using a Nicolet Nexus FTIR spectrometer. The values, given in wave numbers, were rounded to 1 cm⁻¹ using automatic peak assignment. Mass spectra were recorded on a Bruker Esquire 3000 PLUS instrument. The LC textures were studied with a Leica DM4500P POM system that was equipped with a Linkam Scientific Model LTS 350 heating stage and a Canon EOS 60D camera. The melting points were also determined on a Reichert instrument by observing the melting process through an optical microscope. DSC analyses were performed on a Mettler Toledo DSC823e differential scanning calorimeter, using 40 μL aluminum sample pans and Mettler STAR® software for calculation.

Photoresponsive Behavior. The photoresponsivity of the halogen-bonded LCs was studied in planar LC cells (E.H.C. Co., Ltd., alignment layer rubbed polyimide) with a thickness of 2 μm. The temperature was controlled with Linkam Scientific Model LTS 350 heating stage, and the spectra were measured with a Model USB 2000+ spectrometer (Ocean Optics, Inc.) with nonpolarized light from a deuterium–halogen light source. The samples were illuminated with 365 and 395 nm high-power LEDs (Thorlabs, Inc.), equipped with 10 nm, OD 4.0, band-pass filters (Edmund Optics, Ltd.), or with a 457 nm laser. The birefringence (Δn) was measured using a 820 nm laser and crossed polarizers. The sample was placed between the polarizers with a director oriented at angles of ±45°, with respect to the analyzer/polarizer, and the transmitted intensity was measured with a
photodiode. Using the intensity, the birefringence is then defined as follows:

$$\frac{I}{I_0} = \sin^2 \left( \frac{2\pi d n d}{\lambda} \right)$$

The order parameter was determined by measuring the polarized absorption spectra with polarizers positioned parallel ($A_p$) and orthogonal ($A_o$) to the director of the liquid crystal. The order parameter ($S$) was then calculated by determining the absorbances at $\lambda = 420$ nm and using the following equation:

$$S = \frac{A_p - A_o}{A_p + 2A_o}$$

The characterization of the cis-fraction at the onset of the isotropic-to-LC transition (see the Supporting Information for a detailed explanation) is done (i) by determining the trans- and cis-spectra at the isotropic state at 120 °C, and (ii) by assuming the absorbance to be the sum of the trans- and cis-isomers, allowing us to deduce the cis-fraction from the absorption spectra at the onset of the isotropic-to-LC transition. In determining the cis-spectrum, the fact that the absorption spectra are different when measured in dilute dimethyl formamide (DMF), in the LC state, and in the isotropic state, must be noted, and also the thermal isomerization must be taken into account. Because of the thermal isomerization at elevated temperatures (where the samples are isotropic), the photostationary states are measured at different irradiation intensities and the photostationary states are extrapolated to infinite irradiation intensities to remove the effect of thermal isomerization. These photostationary spectra are then used, with the help of Fischer’s method, to determine the cis-spectrum. To deduce the cis-fraction at the second stage, the absorbance of the fully trans-state must be determined to be able to use the deduced spectra of trans- and cis-isomers, and this is done by fitting an exponential curve to the time development of the absorbance. With the spectral information on cis- and trans-states deduced by Fischer’s method, and knowing the absorbance spectra of the fully trans-state through the exponential fitting, the cis-fraction can be determined from the absorbance measurements.

**Synthetic Procedures.** Alkoxystilbazoles are well-known prenecogens, while the iodoperfluorinated azobenzenes were prepared by a NOBF$_4$-mediated diazotization reaction between 4-iodo-2,3,5,6-tetrafluoroaniline and the relative alkoxybenzene. Reactions were carried out in oven-dried glassware under a nitrogen atmosphere. A solution of 4-iodo-2,3,5,6-tetrafluoroaniline (3.715 mmol) in dry acetonitrile (5 mL) was added dropwise into a solution of nitrosonium tetrafluoroborate (3.715 mmol) in dry acetonitrile (5 mL) at −30 °C. After 1 h of additional stirring at −30 °C, a solution of the appropriate alkoxybenzene (7.43 mmol) in dry acetonitrile, was added dropwise. The resulting solution was stirred overnight at room temperature and then water (15 mL) was added. The mixture was extracted 3-fold with CH$_2$Cl$_2$. The organic layers were collected and dried over Na$_2$SO$_4$ and the solvent was removed under reduced pressure. The residue was purified by column chromatography using hexane as eluent to yield the $1\text{-m}$ molecules (20%–25%).

**1-12. Yield, 25%; mp, 67 °C.** 1H NMR (500 MHz, CDCl$_3$): δ = 7.94 (d, J = 8 Hz, 2H), 7.01 (d, J = 8 Hz, 2H), 4.06 (t, J = 6 Hz, 2H), 1.83 (quintet, J = 8 Hz, 2H), 1.48 (quintet, J = 8 Hz, 2H), 1.28 (m, 16H), 0.89 ppm (t, J = 4 Hz, 3H). 13C NMR (126 MHz, CDCl$_3$): δ = 163.68, 147.67, 147.66, 147.61 (ddt, J$_{CF}$ = 265 Hz, J$_{CC}$ = 13 Hz, J$_{CC}$ = 5 Hz), 139.90 (dd, J$_{CC}$ = 265 Hz, J$_{CC}$ = 15 Hz), 133.32 (t, J = 14 Hz, 11.28, 11.05, 7.60 (t, J = 28 Hz), 68.76, 32.08, 29.00–29.65 (4C) 29.51 (2C), 29.27, 26.13, 22.84, 14.25 ppm. 19F NMR (471 MHz, CDCl$_3$): δ = −121.77 (m, 2F), −150.65 ppm (m, 2F). IR: 2919, 2850, 1599, 1578, 1472, 1407, 1252, 1143, 980, 836, 587, 525 cm$^{-1}$. MS/ESI m/z 536.2, found 559.0 (M+Na$^+$).

**X-ray Diffraction.** The azobenzene derivatives and the stilbazoles were separately dissolved in CHCl$_3$ at room temperature in 1:1 ratio, under saturated conditions. The two saturated solutions containing the halogen-bond donor and the halogen-bond acceptor were then mixed in a clear borosilicate glass vial, which was left open. The solvent was allowed to slowly evaporate at room temperature for 3 days until the formation of good-quality single crystals occurred (1-10-2-1, 1-12-2-1, 1-8-2-2).

Single-crystal XRD data were collected on a Bruker AXS KAPPA-APEX II CCD diffractometer using Mo K$_\alpha$ radiation (λ = 0.71073 Å) equipped with Bruker KRYOFOLEX low-temperature device. Data integration and reduction were performed using SaintPlus 6.01. Absorption correction was performed with a multiscan method implemented in SADABS. Space groups were determined using XPREP implemented in APEX II suite. Structures were solved using SHELXS-97 (direct methods) and refined using SHELXL-2008 (full-matrix least-squares on F$^2$) contained in APEX II and WinGX v1.80.01 software packages. All non-hydrogen atoms were refined anisotropically. Hydrogen atoms were placed in geometrically calculated positions and included in the refinement process using a riding model with isotropic thermal parameters. Analysis of crystal data and figures were performed with Mercury 3.5.1.2. Crystal data are reported in Table S1 in the Supporting Information.

PXRD data were collected on a Bruker AXS D8 powder diffractometer was used for all measurements with experimental parameters as follows: Cu K$_\alpha$ radiation (λ = 1.54056 Å), scanning interval, 5°–30° 2θ; step size, 0.016°; and exposure time, 1.5 s/step. Small shifts in 2θ values between the powder patterns of the simulated from single crystal and the experimental XRD spectra are due to different acquisition temperatures. Single crystal structures were collected at 103 or 150 K, while bulk powder materials were recorded at 298 K.

**Computational Analysis.** All geometry minimizations were performed with the M06-2X/def2-TZVP method, using Gaussian 09, Revision D.01. The complexation energy between 2-1 and 1-1 was further obtained with a single-point energy calculation employing the full counter-poise method to correct for the basis set superposition error. An ultra grid was used in all calculations. Default values were used for other parameters. The optimized geometry is given in Figure S1 in the Supporting Information.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemmate.6b03460.

Crystallographic data, (tables, additional pictures, and CIF), powder X-ray data, quantitative analysis of the photoinduced phase transitions, thermal data, computed atomic coordinates (PDF)
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Notes

The authors declare no competing financial interest.

Acknowledgments

P.M. is grateful to the European Research Council (ERC) for funding the Starting Grant FOLDHALO (Grant Agreement No. 307108). The Academy of Finland’s Centres of Excellence Programme (2014–2019) is also acknowledged. A.P. greatly acknowledges the ERC Starting Grant program (Grant Agreement No. 679646), the Academy of Finland Research Fellowship program (Decision Nos. 277091 and 284553), and the Emil Aaltonen Foundation for financial support.

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