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# Effect of halogen substitution on energies and dynamics of reversible photomechanical crystals based on 9-anthracenecarboxylic acid†

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9-Anthracene carboxylic acid derivatives comprise a family of thermally reversible photomechanical molecular crystals. The photomechanical response relies on a [4 + 4] photodimerization followed by dissociation that occurs on timescales of seconds to minutes. A combined theoretical and experimental investigation is undertaken to better understand how chemical modification of the anthracene core influences energetics of both the isolated molecule and the crystal lattice. We use both density functional theory and dispersion-corrected Moller-Plesset perturbation theory computational methods to establish orbital energies, photodimerization reaction energies, and lattice energies for a set of substituted 9-anthracene carboxylic acid molecules. The calculations reveal that steric interactions play a dominant role in the ability to form photodimers and indicate an energetic threshold of 80-90 kJ per mole for the dimerization reaction. Examination of intermolecular bonding in a subset of fluorinated 9ACs revealed the absence of H···F intermolecular bond formation and energy differences that can explain observed trends in the dissociation kinetics and mechanical reset times. Fluorescence recovery after photobleaching experiments shows that the photodimer dissociation kinetics depend on the amount of initial photodimer, preventing a straightforward correlation between halogen atom substitution and dissociation rates using the Bell-Evans-Polanyi principle. The results clarify how molecular structure affects intermolecular interactions and photoreactivity in this family of molecular crystals, but the origin of the complex photodimer dissociation dynamics remains an open question.

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#### Introduction

Crystals composed of photoreactive molecules (photochromes) can transform light energy directly into mechanical work.<sup>1-3</sup> They do this by organizing the molecules that geometrical changes associated photoisomerization, photodimerization, or photocyclization add together in specific directions determined by the crystal axes. Various classes of both intra- and intermolecular reactions have been harnessed to generate this work. In the latter class,  $\begin{bmatrix} 2 + 2 \end{bmatrix}^{4-12}$  and  $\begin{bmatrix} 4 + 4 \end{bmatrix}^{13-16}$  photodimerization reactions are commonly used. To realize thermally reversible (T-type) photomechanical crystals based on

photodimerization, the photodimer must be sufficiently unstable so that its dissociation proceeds rapidly at room temperature. In this context, we have synthesized and studied a class of substituted 9-anthracenecarboxylic acid (9AC) derivatives that crystallize in a head-to-head (hh) geometry due to intermolecular hydrogen bonding between neighboring stacks. The hh geometry gives rise to steric interactions between the COOH groups on cofacial molecules that destabilize the C-C bridge bonds of the photodimer and cause it to spontaneously dissociate back into the monomer pair. This ability of the crystal to reset itself after photodimerization makes it a T-type photomechanical material that could have practical applications.

The 9AC core provides the basis for an entire class of photomechanical molecular crystals because chemical substituents can be added to the anthracene ring in order to tune the material properties. The effects of halogen atom and methyl group substitution on the photodimerization and recovery kinetics have been investigated in two previous papers.<sup>22,23</sup> One surprising result was that F-atom substitution at different carbons around the anthracene ring led to photodimer dissociation rates that varied by more than

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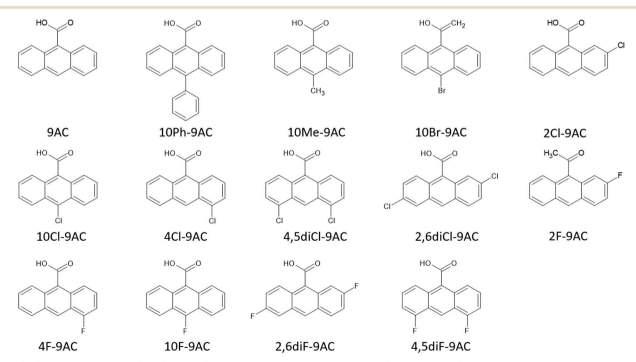
an order of magnitude, despite virtually identical crystal packing geometries. To explain this observation, we hypothesized that intermolecular H···F interactions might affect the relative stabilities of the monomer and photodimer in the different molecular crystals.<sup>23</sup> If the rate of photodimer dissociation can be controlled by the F substitution pattern, this would provide a strategy to design molecular crystals with desired properties, like fast mechanical recovery times.

In this paper, we undertake a combined theoretical and experimental investigation to better understand how chemical modification of the 9AC core influences energetics of both the isolated molecule and the crystal lattice. The goal is to determine the relative importance of local (pairwise) versus nonlocal (crystal) interactions in determining the photochemical reactivity of monomer pairs and the stability of photodimers. We examine a large set of substituted 9AC molecules (Scheme 1) using both density functional theory (DFT) and dispersion-corrected Moller-Plesset perturbation theory (MP2D)<sup>24</sup> computational methods to establish orbital energies, photodimerization reaction energies, and lattice energies. The Bell-Evans-Polanyi (BEP) principle provides a way to correlate the energy difference between a monomer pair and photodimer with the activation energy and thus the thermal dissociation rate.<sup>25–28</sup> These calculations reveal that steric interactions play a dominant role in the ability to form photodimers and suggest there exists an energetic threshold for the formation of the photodimer, but they fail to show any relation between enthalpy differences and observed photodimer dissociation rates. Additional kinetic experiments show that the photodimer dissociation, as measured by the recovery of the monomer fluorescence, is highly nonlinear and depends on the amount of initial photodimer created by the ultraviolet (UV) pulse. The complex reaction kinetics in the crystals prevent a straightforward correlation between halogen atom substitution and dissociation rates. This work clarifies how molecular structure affects intermolecular interactions and photoreactivity in the crystal, but the origin of the complex reaction dynamics in this class of hydrogen-bonded crystals remains unclear.

## Experimental methods

The syntheses of the 10F-9AC, 10Methyl-9AC, 10Cl-9AC, 10Br-9AC, and 10Ph-9AC were previously reported. <sup>22</sup> The syntheses of 2F-9AC, 4F-9AC, and 2,6diF-9AC were described previously. <sup>23</sup> Detailed synthetic procedures for 4,5diF-9AC, 2Cl-9AC, 4Cl-9AC, and 4,5diCl-9AC is described in the ESI.† Crystal structures for all molecules except 4Cl-9AC and 2Cl-9AC are also given in the ESI.†

The measurement of the fluorescence recovery process was performed using an Olympus IX70-inverted microscope. Crystals of the compounds were placed on glass microscope slides. Large rectangular crystals were used for measurements because of their resistance to bending and deformation, preventing misalignment of the pump and probe beams. The alignment of each crystal was confirmed after UV exposure to ensure accurate fluorescence recovery data. Utilizing a pump-probe setup a 405 nm laser was used to irradiate samples for a predetermined time then probed using a less intense 405 nm beam. A Melles Griot Electronic Shutter Controller (04 ISC 850) and a custom LabView program were used to control the pump exposure. The probe spot radius of 17  $\mu m$  was three times smaller than the 55  $\mu m$  pump beam. The beam



Scheme 1 9-Anthracene carboxylic acid derivatives studies in this paper, along with abbreviations use in the text.

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was modulated by a chopper at a frequency of 100 Hz. The resulting signal was obtained using a lock-in amplifier with a photomultiplier tube. To reduce probe beam exposure, intermittent blocking was performed manually during the data collection.

The interpretation of the fluorescence experiments relies on the assumption that the fluorescence is linearly proportional to the monomer population. Previous experiments showed that the fluorescence of the monomer crystal originates from an excimer and is redshifted about 100 nm below the absorption edge,<sup>29</sup> allowing us to assume that reabsorption of the fluorescence by monomer or photodimers is negligible and these optical nonlinearities can be neglected. In the same work, it was found that the loss of fluorescence was linearly proportional to the UV photon dose, again consistent with a linear relation between the loss of monomer and loss of fluorescence. Thus the available data suggests that the fluorescence signal is a good surrogate for the monomer pair population.

### Computational methods

The geometries of gas-phase monomers, dimers, and tetramers of the F-9AC species and their photodimerized analogs were optimized using the BLYP30,31 generalized gradient approximation (GGA) density functional with the D3(BI) dispersion correction<sup>32,33</sup> and the def2-TZVP basis set.34 Frontier orbital energies and quasi-particle gaps were then computed for gas-phase F-9AC monomers using singlepoint energies computed with the range-separated hybrid ωB97X-D density functional<sup>35</sup> in the aug-cc-pVTZ basis set.<sup>36</sup> The highest-occupied molecular orbital (HOMO) energy was equated to the negative of the ionization potential, while the lowest-unoccupied molecular orbital (LUMO) energy was the set to be the negative of the electron affinity. The ionization potentials and electron affinities were computed via the  $\Delta$ SCF approach,<sup>37</sup> taking the energy difference between the neutral species and its corresponding cation or anion. Optical gaps corresponding to vertical excitations were computed via timedependent density functional theory (TDDFT) using the same ωB97X-D/aug-cc-pVTZ model in implicit cyclohexane polarizable continuum solvent.38

The energetics of anthracene photodimerization are notoriously difficult to model, and the results from many common density functionals are both quantitatively and qualitatively incorrect.24,39 Therefore, the F-9AC species and their gas-phase photodimerization reactions in dimer and tetramer clusters were computed with dispersion-corrected second-order Møller-Plesset perturbation theory (MP2D),<sup>24</sup> which been shown to model photodimerization energetics to within a few kcal mol<sup>-1</sup> of coupled cluster theory benchmarks. The MP2D calculations were extrapolated to the complete basis set (CBS) limit by combining Hartree-Fock/aug-cc-pVQZ results with correlation energies extrapolated from the aug-cc-pVTZ and aug-cc-pVQZ basis sets.40 These large basis sets were chosen to help converge the energetics and to reduce basis set superposition error. Because the counterpoise correction<sup>41</sup> is ill-defined for intramolecular interactions, it was not employed here to avoid inconsistencies between the description of the intermolecular  $\pi$ -stacking interactions in the reactants versus the intramolecular interactions in the photodimer product. The TDDFT calculations were performed using Gaussian 16 RevC.01.42 All other gas-phase calculations were performed using PSI4,43 employing density fitting algorithms and the default auxiliary basis sets throughout. The MP2D dispersion correction is available in the current developmental version of PSI4 or can be downloaded as a stand-alone library at https://github.com/Chandemonium/MP2D.

Solid-state calculations on F-9AC crystals were performed using the B86bPBE generalized gradient approximation functional<sup>44,45</sup> and the exchange-hole dipole moment (XDM) dispersion correction<sup>46</sup> using QuantumEspresso 6.4.1,<sup>47,48</sup> a 50 Ry planewave cutoff, and Monkhorst-Pack grids with a k-point density of  $\sim 0.04 \text{ A}^{-1}$ . Core electrons were treated via the projector augmented wave (PAW) approach using PAW potentials for H, C, and F generated with A. Dal Corso's Atomic code v6.1. Experimental crystal structures were relaxed with lattice parameters held fixed at their experimental values. For higher accuracy and consistency with the gas-phase calculations, the single-point crystal lattice energies of the optimized structures were then refined by extracting the key reactive tetramer (two  $\pi$ -stacked pairs of hydrogen bonded dimers), then replacing the DFT energy of that gas-phase tetramer with the corresponding MP2D energy. This approximation is essentially the same as the monomer-correction approach<sup>49</sup> that has proved extremely useful in a number of other organic crystals, 49-51 including the related 9-tertbutyl anthracene ester.<sup>52</sup> The approximation amounts to describing the most important local interactions -those within the key reactive tetramer-with MP2D, while the longer-range contributions arising from the interaction of that tetramer with the surrounding crystal lattice are modeled with periodic DFT. For further insight, the lattice energies were decomposed into contributions arising from within the central tetramer, between the tetramer and the rest of the crystal, and the energy required to distort the monomer from its optimal gas-phase geometry to the crystalline conformation. The interactions in the tetramer were further decomposed into all possible pairwise interactions (consisting of hydrogen bonded pairs,  $\pi$ -stacked pairs, and "diagonal" pairs) plus the residual non-pairwiseadditive "many-body" contribution.

#### Results and discussion

#### A. Effect of halogen substitution on molecular electronic properties

We first wanted to understand how chemical substitution affects the photophysical properties of individual molecules. In the simplest limit, electron withdrawing groups are expected to lower the HOMO and LUMO energies. From a

practical standpoint, this should make the molecule more difficult to oxidize and increase its stability, since reaction with ambient O2 molecules at the C10 and C9 carbons is a major side reaction for photoexcited anthracene molecules leading to 9,10-endoperoxyanthracene derivative that can rearrange to give anthraquinones.<sup>53,54</sup> Fig. 1 shows the calculated gas-phase HOMO/LUMO energy level shifts for the various 9AC derivatives in Scheme 1. As expected, the addition of electron-withdrawing atoms like Cl and F systematically lowers both orbital energies. Surprisingly, Cl substitution seems to have a slightly larger effect, even though it is less electronegative than F. But although the individual orbitals shift by as much as 0.5 eV upon halogen substitution, the HOMO/LUMO shifts tend to offset each other, with the net result that the HOMO/LUMO gap does not change by more than 0.2 eV.

The stability of the HOMO-LUMO gap with respect to chemical substitution is mirrored by the calculated optical gap. Fig. 2a shows representative absorption spectra for 9AC and some fluorinated derivatives. All derivatives retain the characteristic anthracene vibronic lineshape, suggesting that the S<sub>0</sub>-S<sub>1</sub> transition has a similar electronic character in all of them. In Fig. 2b we plot the experimental S<sub>0</sub>-S<sub>1</sub> transition energies in cyclohexane, as well calculated S<sub>0</sub>-S<sub>1</sub> optical energies in implicit cyclohexane solvent for 9AC, 10F-9AC, 2F-9AC, 4F-9AC, 2,6diF-9AC and 4,5diF-9AC. Fluorine substitution causes the measured absorption peak to shift by up to 0.1 eV, while the calculated So-S1 gaps show more scatter but the same magnitude of variation. The calculated So-S1 energies are consistently about 0.1 eV greater than the experimental energies, which is within the typical accuracy for TDDFT calculations.  $^{55,56}$  The experimental and calculated  $S_0$ - $S_1$ 

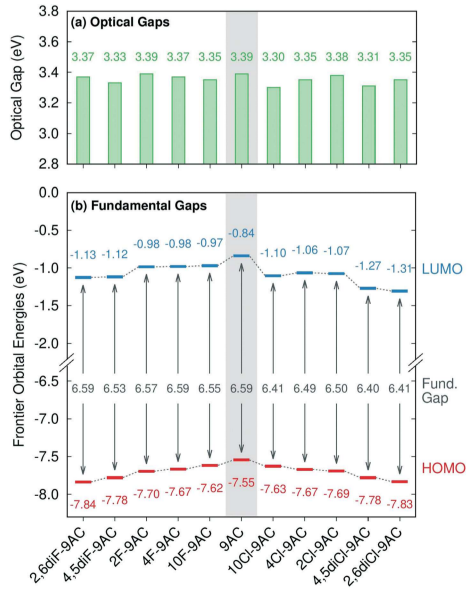
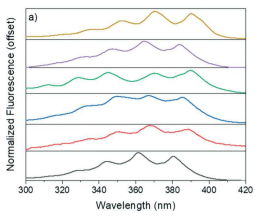


Fig. 1 a) Optical gaps calculated for various 9AC derivatives shown in Scheme 1. b) Effect of halogen substitution pattern on the HOMO/LUMO levels of 9AC.

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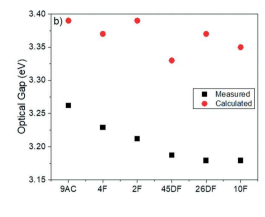


Fig. 2 a) Normalized absorption spectra of 10F-9AC (brown), 4F-9AC (purple), 2,6diF-9AC, 2F-9AC (blue), 4,5diF-9AC (red), and 9AC (black) in cyclohexane b) comparison of calculated (red) and measured (black) optical gaps for fluorinated 9AC compounds

energies and absorption lineshapes are consistent with previous work showing that halogen atom substitution on the anthracene ring can shift the absolute HOMO/LUMO positions but has little effect on the molecule's optical properties. 57,58

#### B. Effect of halogen substitution on dimer energetics

In solution, anthracene carboxylic acids photodimerization typically favors the head-to-tail (ht) conformation. 59,60 In the crystal form, hydrogen bonds between COOH groups on opposing molecules lock the molecules in the hh conformation along the one-dimensional stacks. Gas-phase MP2D calculations comparing the hh and ht arrangements of the photodimer (Table 1) show that the hh photodimer geometry is generally  $\sim$ 32–35 kJ mol<sup>-1</sup> less stable than the ht one due to the steric clash introduced by having the two COOH groups on the same side in the hh geometry. In contrast, the energy differences between the hh and ht geometries of the unreacted gas-phase dimers are only  $\sim$ 3-7 kJ mol<sup>-1</sup>. Taken together, this means that the reaction energies  $\Delta E_{\text{rxn}} = E(\text{photodimer}) - E(\text{unreacted})$  for the hh dimerization are considerably more endothermic than the ht ones (Table 1). This explains why the photodimerization is so easily reversible in the 9AC crystals at room temperature, since only the highly endothermic hh dimerization can occur in

Table 1 Gas-phase MP2D dimer energy differences comparing 9AC derivatives with head-to-head (hh) and head-to-tail (ht) geometries. 1st column: hh versus ht energy difference for unreacted monomer pair; 2nd column: hh versus ht energy difference for dimerized pair; 3rd column: dimerization reaction energy for hh pair; 4th column: dimerization reaction energy for ht pair

Species	Monomer pair $(E_{hh}-E_{ht})$ $(kJ \text{ mol}^{-1})$	Photodimer $(E_{hh}-E_{ht})$ $(kJ mol^{-1})$	$\Delta E_{\rm rxn}(hh)$ (kJ mol <sup>-1</sup> )	$\frac{\Delta E_{\rm rxn}(\rm ht)}{(\rm kJ\ mol^{-1})}$
9AC	4.0	34.3	63.1	32.8
2F-9AC	7.3	34.8	67.0	39.4
4F-9AC	-5.0	35.1	62.3	22.2
10F-9AC	-4.7	-13.5	74.0	82.8
2,6diF-9AC	6.8	32.3	68.1	42.6
4,5diF-9AC	-3.3	32.1	60.4	25.0

the crystalline state. Most of the fluorine substitutions considered impact the energy difference between the ht and hh photodimers minimally (i.e. by 2-3 kJ mol<sup>-1</sup>), except for 10F-9AC where the hh photodimer is actually 14 kJ mol<sup>-1</sup> more stable than the ht photodimer. In this case, the unfavorable electrostatic repulsion between the COOH and/or F groups on the adjacent anthracene units mean that neither the ht nor hh arrangement is particularly favored. Moreover, both hh and ht configurations of 10F-9AC exhibit more endothermic gas-phase photodimerization energies  $\Delta E_{\rm rxn}$  than any of the other derivatives (Table 1), according to these calculations.

MP2D consider the gas-phase photodimerization energies  $\Delta E_{\rm rxn}(hh)$  across a broader set of 9AC derivatives in Table 2, including both ones that have been tested experimentally and other hypothetical monohalogenated 9AC derivatives. The crystal photoreactivity was determined by direct observation of the loss of monomer fluorescence and mechanical changes under 405 nm irradiation. Fig. 3 plots  $\Delta E_{\rm rxn}$  for several 9AC derivatives and indicates whether they exhibit experimental reactivity in the crystal form. The photodimerization of these 9AC derivatives is consistently endothermic, and there seems to be a  $\Delta E_{\rm rxn}$ threshold at around 90 kJ mol<sup>-1</sup>, above which dimer formation prevented. These higher-energy photodimerization reactions have larger substituents located at the 10-position that cannot avoid interacting with each

**Table 2** Gas-phase MP2D predicted dimerization energies ( $\Delta E_{rxn}(hh)$ ) for various **9AC** compounds in the hh configuration, in kJ  $mol^{-1}$ 

Molecule	X = F	X=Cl	X=Br	X=Me	X=Ph
9AC 1X-9AC 2X-9AC 3X-9AC 4X-9AC	$63.1^{a}$ $76.5^{c}$ $67.0^{a}$ $66.5^{c}$ $62.3^{a}$	$ 100.0^{c}$ $69.3^{a}$ $67.9^{c}$ $66.2^{c}$	$ 106.2^{c}$ $72.8^{c}$ $71.7^{c}$ $67.1^{c}$		
10X-9AC 2,6diF-9AC 4,5diF-9AC	$74.0^{a}$ $68.1^{a}$ $60.4^{a}$	$119.9^{b}$ $ 67.8^{a}$	101.7 <sup>b</sup> —	94.9 <sup>b</sup> —	204.7 <sup>b</sup> —

<sup>&</sup>lt;sup>a</sup> Reactive. <sup>b</sup> Non-reactive. <sup>c</sup> Not tested experimentally.

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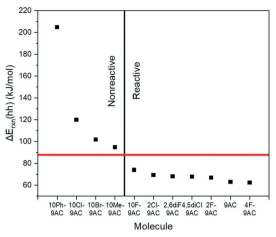
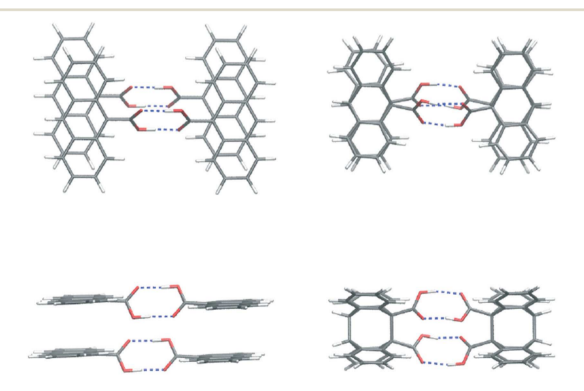


Fig. 3 Predicted gas-phase  $\Delta E_{\rm rxn}$  dimerization energies for various 9AC derivatives, calculated for monomer pairs. The red line represents the approximate energy threshold above which photodimerization is not observed in the crystal form.

other. Among the hypothetical cases, dimerization appears energetically feasible for any fluorine, chlorine, or bromine mono-halogenation at the 2, 3, or 4 positions, with dimerization reaction energies 62-73 kJ mol<sup>-1</sup>. Monofluorination at the 1 position also exhibits a dimerization energy comparable to that of 10F-9AC, while reactions involving larger Cl and Br substituents in the 1 position are unlikely, since they have  $\Delta E_{\text{rxn}}$  of 100 kJ mol<sup>-1</sup> or more. Similar to the 10X-9AC derivatives, these highly endothermic reaction energies stem from steric clashes between the COOH and the halogen groups. It should be noted that  $\Delta E_{\rm rxn}$  is the difference between ground state isomers and thus in principle provides no information about the excited state potential energy surface that determines photodimerization yield. However, we suspect that it indicates the degree of steric interference within the dimer, which is an important factor in determining whether photodimerization proceeds on the excited state. This correlation with sterics probably explains why it correlates with photoreactivity as well.

#### C. Effect of halogen substitution on tetramer and crystal energetics

In order to test our hypothesis that intermolecular bonding plays an important role in the reaction kinetics, we had to examine larger systems. The metastable nature the 9AC photodimer has prevented experimental characterization of its neat crystal structure, although solvates have been investigated.61 Without knowledge of the photodimer crystal structures, the computational modeling instead focused on a gas-phase tetramer cluster model in which two pairs of



#### Non-covalent $\pi$ dimer

#### Covalent Photodimer

Fig. 4 Top and side view of the 9AC tetramer before and after double photodimerization. The photodimer structure was optimized in the gas phase.

**Table 3** Comparison of dimerization reaction energies of fluorinated 9AC compounds in dimer, tetramer with one pair reacted, and tetramer with both pairs reacted ( $\Delta E_{\text{rm}}$ ). The 4th column is an average of the two tetramer reaction energies

Species	Dimer $\Delta E_{\text{rxn}}(\text{hh})$ (kJ mol <sup>-1</sup> )	Tetramer $\Delta E_{\text{rxn}}(\text{hh})$ (kJ mol <sup>-1</sup> )	$\Delta E_{\rm rxn,2}({\rm hh})~({\rm kJ~mol}^{-1})$	$\Delta E_{\rm rxn,avg}(\rm hh) (kJ \ mol^{-1})$
9AC	63.1	73.2	81.7	77.5
2F-9AC	67.0	79.6	83.8	81.7
4F-9AC	62.3	72.1	80.6	76.4
10F-9AC	74.0	84.3	93.4	88.9
2,6diF-9AC	68.1	79.3	88.2	83.8
2,601F-9AC	68.1	/9.3	88.2	83.8

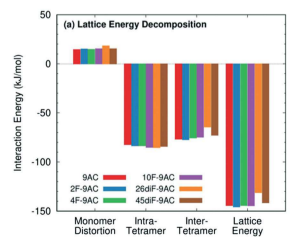
hydrogen bonded monomers could be photodimerized, as illustrated in Fig. 4. These more expensive calculations were performed on a subset of the experimentally investigated molecules in section B. These clusters were extracted from the unreacted monomer crystal, and then either one or both of the monomer pairs in the tetramer were converted to photodimers and the tetramer structure was relaxed. The photodimerization reaction energy in the tetramer is moderately less favorable than in the dimer. As shown in Table 3, the first dimerization within the tetramer is ~11 kJ mol<sup>-1</sup> more endothermic on average, while the second one is  $\sim$ 19 kJ mol<sup>-1</sup> more endothermic. These higher reaction energies reflect the geometric constraints associated with preserving the COOH hydrogen bonds between the stacks. Nevertheless, the qualitative trends in the reaction energies for the different F substitution patterns in the tetramer remain similar to those in the isolated dimer. 4F-9AC again exhibits the lowest  $\Delta E_{\text{rxn}}$ , while **10F-9AC** has largest  $\Delta E_{\text{rxn}}$  and thus would be predicted to be the least stable photodimer.

The similarities of the dimer and tetramer cluster results motivated us to examine the energetics of the monomer crystal lattice in more detail. The goal was to look for any variations in the intermolecular forces that arise from fluorine substitution. The crystal lattice energies are analyzed by decomposing them into contributions arising from (i) distortion of the monomer from its gas-phase to crystalline conformation, (ii) the interaction energy within the key tetramer unit, and (iii) the interaction of that tetramer with the rest of the crystal lattice.

As shown in Fig. 5a, the intermolecular interactions within the tetramer represent the strongest contribution to the lattice energy, but the interactions between the tetramer and the rest of the lattice are almost as large.

One might anticipate the fluorine substitution could create additional hydrogen bonding between tetramer units that would increase the lattice energy. However, the data provides little evidence for the role of intermolecular H···F bonding. With typical H···F distances of 2.3-2.5 Angstroms, any intermolecular H···F bonds are relatively weak here. If such H···F bonds were important, they would impact the interaction energy between the tetramer and the rest of the lattice, but these interactions vary by only a few kJ mol<sup>-1</sup> across most of the derivatives (Fig. 5a). The only notable exception occurs for 2,6-diF-9AC, for which F···F repulsion actually weakens these interactions by about ~10 kJ mol<sup>-1</sup> relative to the other derivatives. Fluorine substitution has only minor impacts on the other energy contributions as well. For example, the interactions within the tetramer vary by only  $\sim 2$  kJ mol<sup>-1</sup> across the derivatives, and the differences among the monomer distortion energies are similarly small (with 2,6-diF-9AC again being a moderate exception). In the end, the total lattice energies for all derivatives lie in the 141-146 kJ mol<sup>-1</sup> range, except for 2,6diF-9AC at 131 kJ mol-1 (due to the aforementioned F···F repulsion and larger distortion energy penalties).

It seemed surprising the F substitution had such small impacts on the tetramer energies, since it is known that



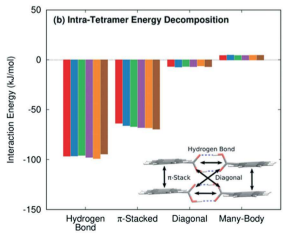


Fig. 5 a) Effect of fluorine substitution on various lattice energy terms for six different 9AC derivatives. b) Effect of fluorine substation on tetramer energy terms for the same 9AC derivatives. In all cases, the changes induced by different locations of the F-atom amount to only a few kJ mol<sup>-1</sup>.

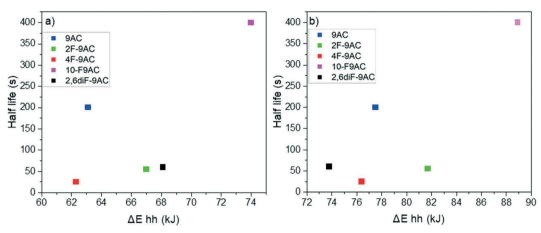


Fig. 6 Fluorescence recovery times  $(\tau_{1/2})$  taken from ref. 23 plotted versus the calculated a) dimer and b) tetramer photodimerization energies.

halogenation impact pi-stacking interactions can appreciably.<sup>62</sup> Fig. 5b further decomposes the intermolecular interactions within the key tetramer into the sum of the two pairwise hydrogen bond interactions, the sum of the two  $\pi$ -stacking energies, the two pairs of "diagonal" interactions, and the residual many-body contribution arising from the beyond-pairwise interactions within the tetramer. As expected, we do observe that fluorination systematically increases the strength of the  $\pi$ -stacking interactions by up to 5 kJ mol<sup>-1</sup>, and the di-fluorinated species stacks are stabilized more than the mono-fluorinated ones. However, the other interactions vary less systematically with fluorine substitution, and the net effect is that the improved  $\pi$ -stacking energetics are largely cancelled out in the tetramer as a whole. This conclusion is consistent with the fact that most of the derivatives exhibit almost identical crystal packing (except for 10F-9AC). Overall, these results highlight how the details of crystal packing stability on the whole can defy simple intuitive logic derived from knowledge of individual pairwise intermolecular interactions.

Given the weak effect of F-substitution on the crystal energetics, it is perhaps not surprising that we find no strong correlation between the energetics and the dimer dissociation rate. According to the BEP principle,  $\Delta E_{\rm rxn}$ should also correlate with the dissociation rate through its activation energy. In Fig. 6a, we plot the dissociation half lives ( $\tau_{1/2}$  from ref. 23) versus the dimer  $\Delta E_{\rm rxn}$  for molecules 9AC, 10F-9AC, 2F-9AC, 4F-9AC, and 2,6diF-9AC. Unlike Fig. 3, there is no obvious correlation. The inclusion of dimer-dimer interactions by using a tetramer cluster does not change this situation, as shown in Fig. 6b. In both cases, the largest predicted hh photodimerization energy  $\Delta E_{\rm rxn}$  in Table 1 is found for 10F-9AC, while the smallest one occurs for 4F-9AC. If the dimer → monomer pair reactions of the different derivatives have similar transition state structures, the BEP principle would predict the 10F-9AC dimer to have the fastest dissociation rate and 4F-9AC to have the slowest rate. This is precisely the opposite of what is observed experimentally.

#### D. Nonlinear photodimer dissociation kinetics

The absence of a correlation between  $\Delta E_{\text{rxn}}$  and the photodimer lifetimes in Fig. 6 suggests that the BEP principle cannot be naively applied to this class of reactions. One assumption of the BEP model is that the Arrhenius prefactors and transition state structures are similar within the family of compounds under study. While this assumption seems reasonable for the 9ACs, it may be that the transition state is much more sensitive to the surrounding crystal environment than the minimum energy monomer and dimer configurations. A second possibility is that the accurate calculation of the photodimer energies requires taking the full crystal lattice into account, rather than isolated clusters. However, there is no hint from the monomer calculations that the inclusion of such terms would shift the relative energies by more than a few kJ mol<sup>-1</sup>. Rather than focusing on energetics, a third explanation centers on the possibility that the bulk photodimer dissociation kinetics are more complicated than the simple two-state model that underlies the BEP principle.

In the ideal case, the photodimerization can be described by a simple two-state model subject to well-defined forward and backward first-order rate constants that are independent of concentration. This assumption is usually valid in dilute solution, where the molecules react independently in a homogeneous environment. But in a crystal, where the reacting molecules are in intimate contact, the surrounding can change as the fraction of dimers evolves, leading to a time-dependent rate that changes as the reaction progresses. 63-65 In this limit, a single rate cannot describe the process and it is no longer possible to make a straightforward connection between equilibrium energies and dynamics. One hallmark of such nonlinear kinetics is their dependence on initial concentrations. Recent studies on the kinetics of the dimer dissociation in 4F-9AC crystals showed strong deviations from first-order exponential behavior, where a larger initial photodimer concentration slowed down the dissociation reaction and the recovery of CrystEngComm

the monomer fluorescence signal.<sup>29</sup> The recovery was also sensitive to the presence of a 100× weaker probe beam that could stall the monomer recovery. We interpreted these concentration-dependent decays as evidence of cooperative dynamics, perhaps related to nucleation.

To determine whether the linear kinetic model is valid for the other 9AC derivatives, we measured the kinetics of monomer recovery using a fluorescence recovery after

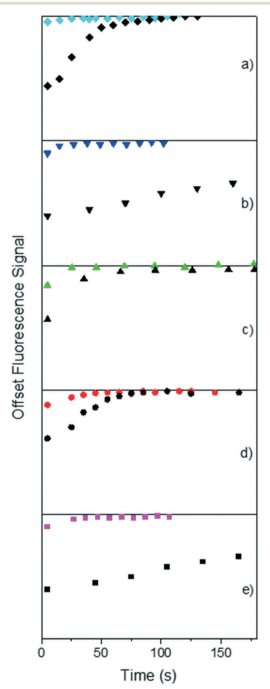


Fig. 7 Fluorescence recovery after photobleaching curves for a) 4F-9AC, b) 10F-9AC, c) 2,6diF-9AC, d) 2F-9AC, and e) 9AC after small (~5% colored symbols) and large (~50% black symbols) photodimer conversions. The data in a) is taken from ref. 29.

photobleaching (FRAP) experiment. We were careful to only use an intermittent probe technique so the fluorescence detection did not perturb the recovery and found concentration-dependent kinetics in all of them to varying degrees. Fig. 7 compares the high and low bleach (initial photodimer concentration) recovery curves for molecules 9AC, 10F-9AC, 2F-9AC, 4F-9AC, and 2,6diF-9AC. All exhibit slower decays for large bleach depths (high initial photodimer conversion), with the effect being most pronounced for 9AC and least pronounced for 2,6diF-9AC and 10F-9AC. Note that 9AC can be grown in two different polymorphs, monoclinic and triclinic.<sup>22</sup> This different symmetry arises from the slightly different arrangement of the one-dimensional stacks along the b-axis, as detailed in the ESI.† The same nonlinear behavior was seen in both polymorphs (ESI†), providing additional experimental evidence that subtle changes in stack orientation do not strongly affect the dissociation kinetics.

As in our previous paper,23 we can parameterize the data in Fig. 7 by extracting  $\tau_{1/2}$ , the time required for the fluorescence recovery to achieve 50% of its pre-bleach value. These values are summarized in Table 4. For low bleach depths (low initial photodimer concentrations), the  $\tau_{1/2}$ recovery times are similar for all molecules except 10F-9AC, as shown in Fig. 8a. For deeper bleaches, Fig. 8b shows how the  $\tau_{1/2}$  times diverge by a factor of 10. From Fig. 8a, it appears that any discrepancy between the  $\tau_{1/2}$  values for different compounds is magnified when large (>20%) photodimer conversions are measured. The similarity of the low conversion recovery rates among the different compounds suggests that local energetic considerations, predicted by theory to be similar, are controlling the dissociation of dilute dimers. At higher concentrations, however, it is possible that many-body effects become important, and these cannot be identified by the dimer and tetramer level of theory employed here. We note that 10F-9AC continues to be an outlier, even for low conversions.

In an earlier paper, we did not realize that the bleach depth and the presence of the probe beam could have a significant effect on the recovery dynamics.<sup>23</sup> Although the qualitative conclusions from that work are still valid, the quantitative comparison of recovery times is considerably more complex than we appreciated at that time. The origin of the variable dissociation behavior among the 9AC derivatives remains a mystery. The kinetic behavior of a photodimer surrounded by

Table 4 The half times from shallow ( $\sim$ 10%) and deep ( $\sim$ 50%) fluorescence bleaches (fraction of crystal converted to the nonfluorescent photodimer) for selected 9AC compounds

	Shallow bleach recovery $(\tau_{1/2})$ (s)	Deep bleach recovery $(\tau_{1/2})$ (s)
9AC	7.2	174.5
2F-9AC	15.3	54.4
4F-9AC	6.5	31.5
10F-9AC	270.1	333.2
2,6diF-9AC	6.8	21.5

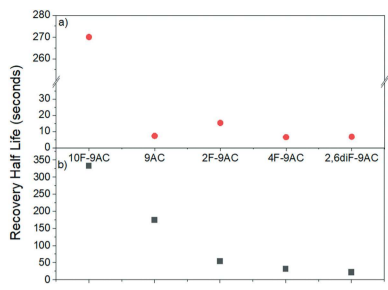


Fig. 8 Fluorescence recovery times ( $\tau_{1/2}$ ) for 9AC, 10F-9AC, 4F-9AC, and 2,6diF-9AC compounds after a) small ( $\sim$ 5%) photodimer conversion and b) large ( $\sim$ 50%) photodimer conversion.

monomer pairs must be quite different from that of one surrounded by other photodimers. It is possible that the strong intermolecular hydrogen-bond interactions in the 9AC family make these compounds more susceptible to such cooperative phenomena. It is also possible that this kinetic phenomenon is affected by presence of crystal defects, which could vary between derivatives. One clue to the origin of the nonlinear kinetics is that 2,6-diF-9AC appears to be the least susceptible to the cooperative effects, with the recovery times at both high and low bleach depths being quite similar. This relative insensitivity is consistent with its weaker intermolecular interactions as predicted by calculations shown in Fig. 5a. Weaker interactions can allow the dissociation kinetics to be more effectively decoupled from neighboring sites and lead to more standard first-order kinetics. Even though a direct connection to the rate is not possible because there is no single, well-defined Arrhenius rate, this observation provides a starting point for future investigations.

#### Conclusion

To summarize, extensive calculations of the crystal energetics and kinetic measurements for a series of 9AC derivatives have provided insight into the reaction dynamics of this class of T-type photomechanical crystals. The important role of steric interactions and hh packing in determining the ability to create a metastable photodimer in the crystal was clarified. examination confirmed and Careful intermolecular bonding in a subset of fluorinated 9ACs revealed an absence of H···F intermolecular bond formation and a lack of clear energy differences that could explain observed trends in the dissociation kinetics and mechanical reset times. Instead, it appears that differences between molecules reflect variations in cooperative phenomena that give rise to highly nonlinear (concentration dependent) kinetics. It is possible that such kinetics are especially pronounced in the **9AC** series due to the presence of strong intermolecular COOH bonds. With a more detailed understanding, it may be possible to design molecular structures that optimize intermolecular interactions to amplify or suppress such collective phenomena and create light-responsive crystals with novel properties.

#### Conflicts of interest

There are no conflicts to declare.

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