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Introduction

Over the past decades, molecular switches have become ubiquitous in the development of smart materials, rendering reversible changes in material function upon input stimuli.^{1,2} In addition to designing new molecular switches and the synthetic optimisation of existing structures, the deeper understanding of their fundamental responsiveness has led to a vast body of research on small molecular photoswitches, most notably on azobenzenes,^{3,4} spiropyrans^{5,6} and diarylethenes.⁷ These exemplary molecular photoswitches have proven to be excellent building blocks for integration in materials owing in particular to their multi-responsiveness, reacting not only to light,^{8,9} but also, *e.g.*, heat,¹⁰ mechanical input,¹¹ and pH-changes.¹² It is particularly attractive when there is synergy between distinct stimuli that leads to new functionality, as is the case for the photo- and pH-response in spiropyrans, where weak acids slow down the thermal reversion and, as more recently shown, strong acids can open up to new photophysics in the protonated isomers.^{6,13} Conversely, for the azobenzene photoswitch, the interaction with acids has been long shown to catalyse rather than retard the reversion of *Z*-to *E*-isomer

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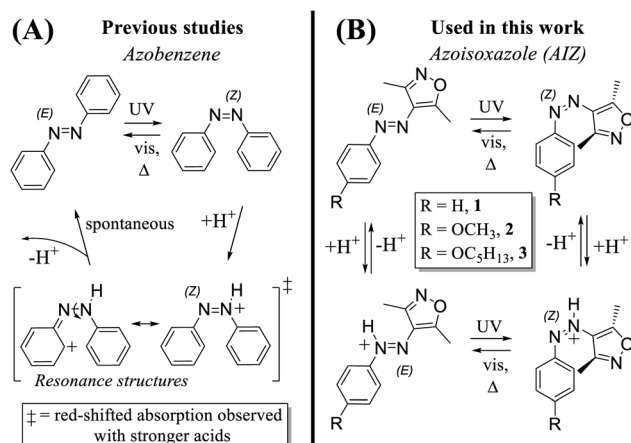
Acid-catalysed liquid-to-solid transitioning of arylazoisoxazole photoswitches†

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Molecular photoswitches play a vital role in the development of responsive materials. These molecular building blocks are particularly attractive when multiple stimuli can be combined to bring about physical changes, sometimes leading to unexpected properties and functions. The arylazoisoxazole molecular switch was recently shown to be capable of efficient photoreversible solid-to-liquid phase transitions with application in photoswitchable surface adhesion. Here, we show that the arylazoisoxazole forms thermally stable and photoisomerisable protonated *Z*- and *E*-isomers in an apolar aprotic solvent when the pK_a of the applied acid is sufficiently low. The tuning of isomerisation kinetics from days to seconds by the pK_a of the acid not only opens up new reactivity in solution, but also the solid-state photoswitching of azoisoxazoles can be efficiently reversed with selected acid vapours, enabling acid-gated photoswitchable surface adhesion.

(Scheme 1).^{14–20} The extent of catalytic reversion depends on the exact acid strength and the amount added, as well as the solvent type in conjunction with its limiting pK_a .²¹ *I.e.*, in 1938 Hartley already reported that addition of several drops of acetic acid to water accelerates the optical *Z*-to *E*-conversion to several seconds,¹⁴ while Ciccone *et al.* found no effect upon addition of acetic acid to *Z*-azobenzene in aqueous ethanol.¹⁵

Acid-catalysed reversion rates in azobenzenes were reported over the years to be independent of ionic strength of the



Scheme 1 (A) Acid-catalysed *Z*-to *E*-reversion reported for azobenzenes. The isomerisation occurs spontaneously, a lingering red-shifted azonium absorption is observed when strong acids are used. (B) The (pH-gated) photochromism and acidochromism of arylazoisoxazole reported here. We note that the depicted protonation sites are based solely on an extensive theoretical analysis (see below).



acids,^{15,16} with the isomerisation step of the *Z*-to *E*-cation determined as the rate determining step.¹⁷ The accelerated isomerisation is commonly attributed to the mesomerism of the azonium ion weakening the azo double bond,^{14–17,19} an effect that was observed to be stronger when electron-rich substituents are conjugated to the azo double bond.^{14,16} Interestingly, despite the strong pH dependence of the *Z*-isomer, the absorption of the plain *E*-azobenzene was found to be independent of aqueous HCl between pH 1 and 13.¹⁶ Indeed, the formation of azonium species in some (*para*-) substituted azobenzenes was observed to require much harsher conditions, such as a 6 M HClO₄ solution,^{17,20} or gas phase ionisation.²²

Alternatively, aminoazobenzenes, such as the methyl orange pH-indicator, have been shown to yield red-shifted azonium ions on account of resonance contributions from the amino-substituent,¹⁹ with 4,4'-diaminoazobenzenes even protonating up to pH 4.²³ Such red-shifting of absorption through azonium formation runs not only parallel to the strategy of appending auxiliary functional groups,^{24–27} it also gives rise to new isomerisation kinetics altogether. The group of Woolley, for example, discovered that the *pK_a* of the azo-moiety is significantly increased owing to hydrogen-bonding in tetra-*ortho*-methoxy azobenzenes, such that the protonated species is preferred at physiological conditions.^{28–31} Their red-shifted absorption opens up to *E*-to *Z*-isomerisation with NIR light with lifetimes in the order of milliseconds to seconds,^{28–30} leading to application in various smart materials.^{32–35} Importantly, the tetra-*ortho* substitution was found to be vital in raising their *pK_a* to such extent, while a symmetric shift of two methoxy units to the *meta*-position caused a *pK_a* drop of 5 units.²⁹

Despite the rapidly grown interest in photochromically superior azoheterocycles as an alternative to azobenzenes,³⁶ only few contributions have been concerned with the generation and stabilisation of their azonium species.^{37–39} For example, Fuchter *et al.* showed that for azopyrazole^{40–42} (AAP) the acid-catalysis of azobenzenes is retained when trichloroacetic (CCl₃COOH), trifluoroacetic (CF₃COOH) and hydrochloric acid (HCl) are applied, and that the transient *Z*-azonium AAP could even be observed during catalysis.³⁷ In further work, *ortho*-substitution of the azopyrazole benzene ring analogous to the work of Woolley *et al.* was found to have an effect on the lifetime of *Z*-azopyrazoles also, albeit without mention of its effects on acid-catalysis.³⁸ More recently, Walther *et al.* succeeded in demonstrating the general acid-catalysis of water-soluble azopyrazoles, opening up to a tunability of *Z*-isomer half-lives over 5 orders of magnitude depending on the pH value.³⁹ This retention of the acid catalysis in the AAP azoheterocycle brings about important implications for azoheterocycles in general, as their superior photophysics, and their vastly increased thermal half-lives in particular, give rise to a significantly larger control over half-life times without the need for synthetic adjustments.

Azoisoxazoles (AIZs), as a relatively new type of azoheterocycle, show great promise as components in smart materials owing to the photoinduced reversible solid-to-liquid phase transition demonstrated recently.⁴³ Due to the sterically

induced out-of-plane bending of the *Z*-isomer the intermolecular interactions are incapacitated, leading to a drastic decrease in melting point and consequent liquefaction of the azoisoxazole upon UV-induced isomerisation. Previously, we have utilised this remarkable retention of photophysics in the solid state to develop photoreversible low molecular weight (LMW) adhesives, cohering glass surfaces at 30.8 N cm^{−2} with rapid debonding upon UV-irradiation.⁴⁴ Although the control over the bonding/debonding process is remarkably swift and reversible even upon physical breaking of the cohesion, the photochemical approach requires the use of at least one transparent surface. In this contribution, we seek to expand on the promising reusable photophysical features of these low molecular weight azoisoxazoles by exploring the possibility to kinetically control their solidification with protons. Though we demonstrate that the acid-induced isomerisation takes place with select acids for azoisoxazoles also, the terms “weak acid” and “strong acid” turn out to be imperative while use of acids stronger than the conventional HCl and CF₃COOH, *i.e.*, trifluoromethanesulfonic acid (CF₃SO₃H, *pK_a* = 0.7 in acetonitrile⁴⁵), yield steady state *Z*-azonium (*Z*-AIZH⁺) and *E*-azonium (*E*-AIZH⁺) species as observed by UV/vis and ¹H NMR spectroscopy. Furthermore, the isomerisation kinetics are not only *pK_a*-tunable in solution, but also in the pristine liquefied *Z*-form by exposure to acid vapours. Ultimately, it empowers the versatile adhesive with a rapid, non-destructive trigger for reversibly conjoining surfaces with remarkable adhesive strengths irrespective of transparency, and showcases the further potential that the added pH-gated control opens up to for this already widely versatile class of photochromes.

Results and discussion

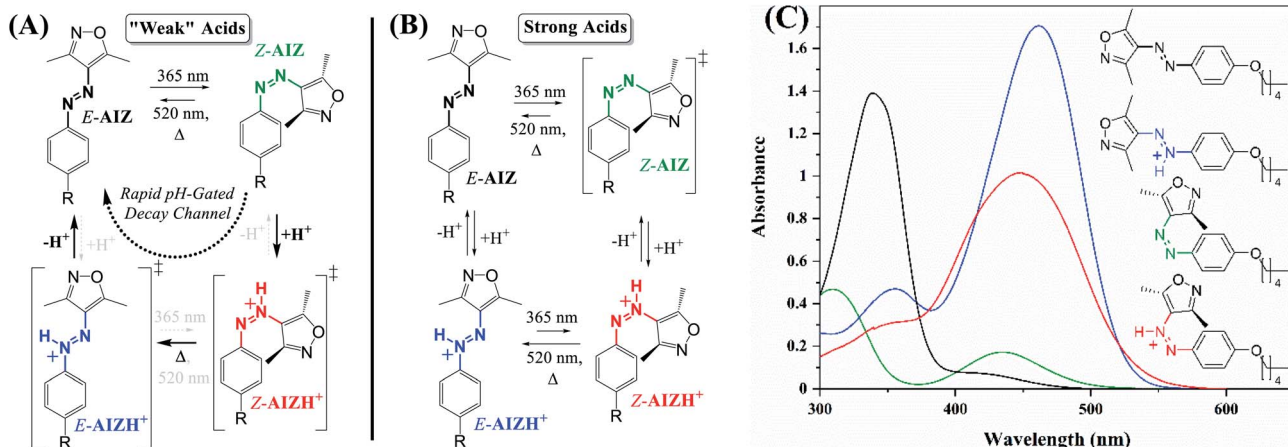
Azonium formation in azoheterocycles

Contrary to the catalysed azo-reversion with conventional acids such as CF₃COOH and HCl, we observe that the addition of the more acidic trifluoromethanesulfonic acid (CF₃SO₃H) to azoisoxazole AIZ 1 causes a shift to a new steady-state absorption (Fig. S1†). Comparably, the analogous unsubstituted AAP 4 (ref. 41) also shows a steady-state absorption upon addition of CF₃SO₃H (Fig. S2†) and, furthermore, distinct azonium chemical shifts in ¹H NMR spectroscopy (Fig. S3,† for AIZ spectra see below, “*Thermally stable Z-and E-protonation with strong acids*”). Although the addition of CF₃SO₃H to the unsubstituted AIZ 1 yields a new absorption at 315 nm, the alkoxy AIZ 2 and 3 azonium ions (AIZH⁺) exhibit strong red-shifted absorptions instead (Fig. 1), while undergoing catalytic *Z*-to *E*-reversion in response to HCl and CF₃COOH. Hence, it is possible to capitalise on varying responses and response times by judicious choice and quantity of acid, opening up to a wider range of control over previously established as well as new physical properties.

Accelerated *Z*-to *E*-reversion with weak acids

Owing to mesomerism,⁴⁶ the uncatalysed half-life of the 4-alkoxy-substituted *Z*-AIZs 2 and 3 in solution was determined to





be 7 d (ref. 44) rather than the 45 d for Z-AlIZ 1.⁴³ Although we observe no reactivity of Z-AlIZ 2 and 3 to a large excess of acetic acid (CH_3COOH , 5000 equivalents in acetonitrile, Fig. S8†), addition of CF_3COOH and HCl cause the half-life to diminish to the (sub)second timescale at near-stoichiometric amounts already (Fig. S9 and S10†). DFT calculations of activation free energies confirm that the Z-to E-barrier is substantially reduced in the azonium species from 1.29 eV (versus 1.11 eV experimentally⁴⁴) to 0.86 eV (Fig. 2A). Importantly, out of three possible N-protonation sites, the azo-N at the phenyl ring is lowest in energy for the E-form, and the N-azo at the heterocycle for the Z-form (Fig. S11,† the protonated heterocycle being highest in both cases; we note that our calculations on AAP 4 are in line with those by Fuchter *et al.*,³⁷ Fig. S12†). The greatly reduced thermal Z/E barrier accessible at excess HCl or

CF_3COOH causes the generated Z-AlIZH^+ isomer to rapidly equilibrate with E-AlIZH^+ . However, although the pK_a of HCl is sufficient to interact with the Z-form, any E-AlIZH^+ generated is spontaneously deprotonated due to its significantly increased acidity. The thermodynamically trapped unprotonated E-AlIZ (barrier of 1.86 eV to Z-AlIZ) that results from this energetically favourable protonation–isomerisation–deprotonation sequence is thus continuously and, moreover, rapidly formed owing to Le Chatelier's principle (Fig. 1A; further supported by calculations taking into account conjugate bases, Fig. S13†).

Notably, catalytic amounts of HCl or CF_3COOH cause the reversion to occur gradually, allowing for the thermal reversion to be moderated as described for azobenzenes and azopyrazoles before (Fig. 2B, S14†).^{14–20,37,39} Though the tuneable half-life ranges from several seconds at stoichiometric equivalents of

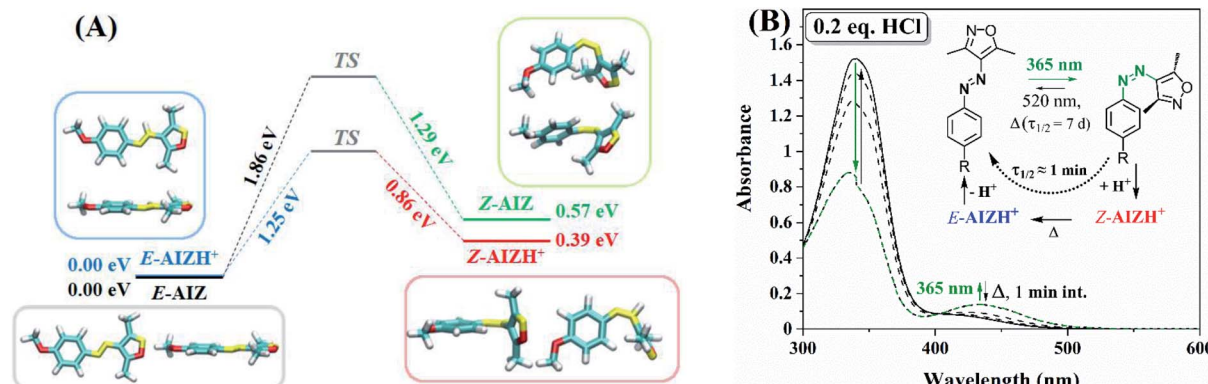


Fig. 2 (A) Relative Gibbs free energies of the unprotonated and protonated isomers of AlIZ 2 and their transition states. For details on the DFT calculations see the experimental section. (B) UV/vis absorption spectra of AlIZ 3 (70 μM in acetonitrile) with 0.05 eq. of HCl (black solid line). The thermal reversion is sped up through the acid-catalyzed reversion on the right, hence the PSS_{365nm} (radiant flux 1.2 W) is compromised through spontaneous reversal via the azonium ions (black dashed lines, spectra every 2 min). Note also that the isosbestic point at 403 nm, where the molar absorptivity of both azonium species is significantly higher (Fig. 1C), rules out a steady state generation of the Z-AlIZH⁺ isomer.

HCl to minutes at catalytic amounts, the higher pK_a and solubility of trifluoroacetic acid allow for more control at superstoichiometric quantities (Fig. S15†), yielding varying well-tunable half-lives (Table 1). The reduction in half-life eventually stagnates upon increasing the acid content, analogous to the earlier reports on the deviation of first-order kinetics in azobenzenes.^{14,15,17,37,39} We note, however, that this is likely due to the weak but increasing involvement of the *E*-azonium isomer, not least because an increase to 5000 eq. CF₃COOH causes the half-life to significantly rise again. In other words,

Table 1 Thermal half-life of Z-AIZ 3 (70 μ M in acetonitrile) at varying concentrations of CF₃COOH as determined by UV/vis spectroscopy

CF ₃ COOH	Thermal half-life	Rate constant
0 eq. (ref. 44)	7 d	—
2 eq.	338.4 s	± 46.3
10 eq.	242.1 s	± 11.1
50 eq.	118.8 s	± 18.9
200 eq.	43.5 s	± 8.2
1000 eq.	24 s	± 1.2
2000 eq.	25.2 s	± 0.5
5000 eq.	59.9 s	± 3.5

despite the mismatch in pK_a the *E*-AIZH⁺ becomes energetically more feasible in presence of thousands of possible (transient) proton sources, increasing its life-time and thus delaying the catalytic process.

Theoretical analysis of the Z \rightarrow E thermal conversion

The large change in thermal reversion rates upon protonation was investigated in further detail through locating the accessible regime of the corresponding potential energy hypersurfaces employing DFT. For azobenzenes, thermal reversion pathways have been subject to many theoretical and experimental studies.^{4,47–50} The molecular motion during azoisomerisation in general is typically described as comprising rotation and inversion, and certain combinations thereof (Scheme S1†).⁴ Classically, quantum chemical calculations rely on locating individual stationary points (minima, transition states) and lack information on how these points are dynamically connected. Here, we adopt a point of view more suited to a dynamical interpretation and scan the potential energy surface. Fig. 3 presents relaxed energy profiles starting out of the minimum energy valley of AIZ 2 at varying CNNC dihedral angles and C(5-ring)NN bond angles.

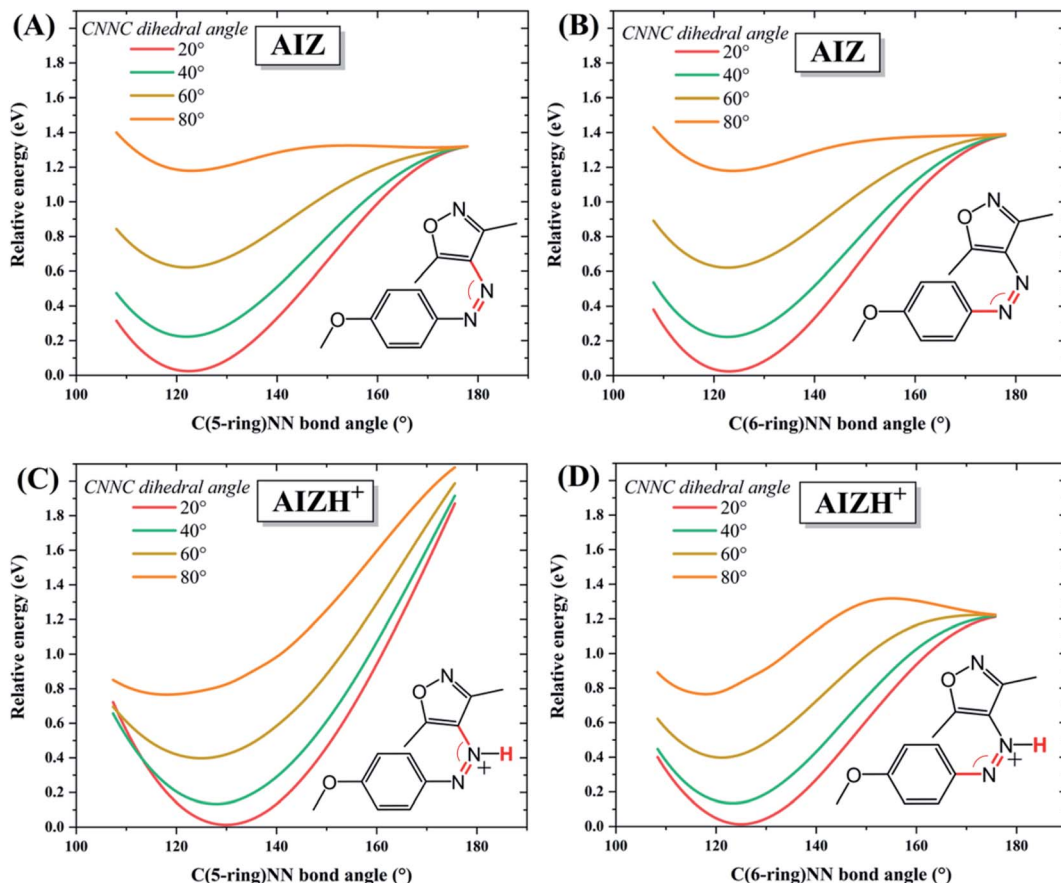


Fig. 3 Relative potential energies of (A and B) AIZ 2 and (C and D) AIZH⁺ 2 along the coordinates of the dihedral angle CNNC (fixed at the values specified), and either (A and C) C(5-ring)NN bond angle or (B and D) C(6-ring)NN bond angle. The amount of data was reduced here for better visualisation, additional data at 10° intervals of CNNC dihedral angle is reported in Fig. S16†.



Accordingly, for the neutral as well as the protonated AIZ species, the minimum energy pathway is dominated by an increase of the dihedral angle CNNC (commonly viewed as 'rotation') up to 80° whereupon an opening of the bond angle CNN becomes feasible. While the neutral AIZ allows bond angle opening to both sides, for the protonated AIZ opening at the non-protonated side is feasible only. Hence, the mechanism involved is labelled 'rotation-assisted inversion' for the neutral and '(pure) rotation' for the protonated AIZ species (Scheme S1†). The corresponding results for the (neutral isomers of) AAP 4 and the parent compound azobenzene are effectively the same as for the unprotonated AIZ 2 (Fig. S17†), and thus constitute a general feature of the azo moiety, in line with recent experimental⁴⁹ and theoretical⁵⁰ reports. Protonation of Z-AIZ can occur at either of the two azo-nitrogens, adjacency to the 5-ring being energetically favoured by less than 0.1 eV (Fig. S11†). Additionally, the likelihood of protonation *via* the solvent, investigated by minimum energy path calculations for proton transfer from either azo-nitrogen to a single explicit molecule of acetonitrile, CH_3COO^- , CF_3COO^- , or CF_3SO_3^- (Fig. S13†), further supports our experimental observations. The acidity of CH_3COOH is far too weak to protonate AIZ, and while for CF_3COOH protonation of AIZ is considerably more likely, the proton still slightly favours the conjugate base of the acid. In the case of $\text{CF}_3\text{SO}_3\text{H}$, however, the equilibrium strongly shifts to AIZ. Concerning a possible shift of the proton from one azo-nitrogen to the other, Fig. S13† reveals that a frequent exchange should take place *via* CF_3COO^- due to a very shallow potential landscape for both isomers, whereas for CF_3SO_3^- it is only feasible for the *E*-isomer, though less frequent than for CF_3COO^- .

Thermally stable Z-and *E*-protonation with strong acids

Although substitution patterns play a vital role in increasing the pK_a of *E*-azonium towards physiological conditions,^{19,23,28–30} the access to a larger range of molecular properties in less complex existing azo-structures is highly advantageous. To achieve this, the pK_a of the acid needs to be near the low pK_a of the azo-bond.

Whereas neither of the azonium isomers are observed by UV/vis spectroscopy even at 500 equivalents of HCl, addition of increasing amounts of $\text{CF}_3\text{SO}_3\text{H}$ to *E*-AIZ 3 causes a stepwise formation of the *E*-AIZH⁺ azonium with saturation at about 1500 equivalents (Fig. S18†). Accordingly, by considering the equilibrium being practically exclusively between *E*-AIZ and *E*-AIZH⁺ and assuming full dissociation of $\text{CF}_3\text{SO}_3\text{H}$, the pK_a of the *E*-isomer is approximated to be -2.48 ± 0.16 by UV/vis spectroscopy.

Interestingly, a sharp cut-off is observed for protonation in ¹H NMR spectroscopy instead, transitioning from unprotonated to protonated between 5 and 6 equivalents of $\text{CF}_3\text{SO}_3\text{H}$ (Fig. S19†). We note, however, that adding multiple equivalents of $\text{CF}_3\text{SO}_3\text{H}$ to 3.3 mM AIZ in ¹H NMR spectroscopy (*versus* 70 μM in UV/vis) results in a much lower pH and thus increased possibilities of interaction. This observation was made by the group of Fuchter also while protonating an AAP with HCl, requiring several orders of magnitude less equivalents at ¹H NMR concentration.³⁷ Presumably, the high interaction is also the reason for the sharper cut-off in chemical speciation than in UV/vis absorbance spectroscopy.

Alternatively, addition of $\text{CF}_3\text{SO}_3\text{H}$ at the PSS_{365nm} (97% *Z*-isomer) yields an additional red-shifted absorption in UV/vis spectroscopy (Fig. S20†) as well as distinct *Z*-AIZH⁺ chemical shifts in ¹H NMR spectroscopy (Fig. 4, S21, Table S1†). Importantly, neither the *Z*-AIZH⁺ nor *E*-AIZH⁺ chemical shifts are observed upon addition of equal amounts of $\text{CF}_3\text{SO}_3\text{Li}$ to AIZ 3 (Fig. S22†). Furthermore, whereas near-equivalent amounts of $\text{CF}_3\text{SO}_3\text{H}$ destabilise the *Z*-AIZ already, its presence is unaltered with 15 eq. of $\text{CF}_3\text{SO}_3\text{Li}$, excluding a significant role of ionic strength on chemical speciation. The *Z*-AIZH⁺ species generated in presence of $\text{CF}_3\text{SO}_3\text{H}$, unlike the unprotonated *Z*-form, is metastable at room temperature and reverts to a thermodynamic equilibrium with 94% *E*-AIZH⁺ over time. The larger thermodynamic ratio of the *Z*-to *E*-azonium in presence of $\text{CF}_3\text{SO}_3\text{H}$ in ¹H NMR spectroscopy (6% *Z*-form as opposed to *ca.* 1% in absence of a proton source⁴⁴) is in line with the change in energetic barriers observed by DFT (Fig. 2A). Moreover, only 2 new sets of signals are observed here, their kinetics being

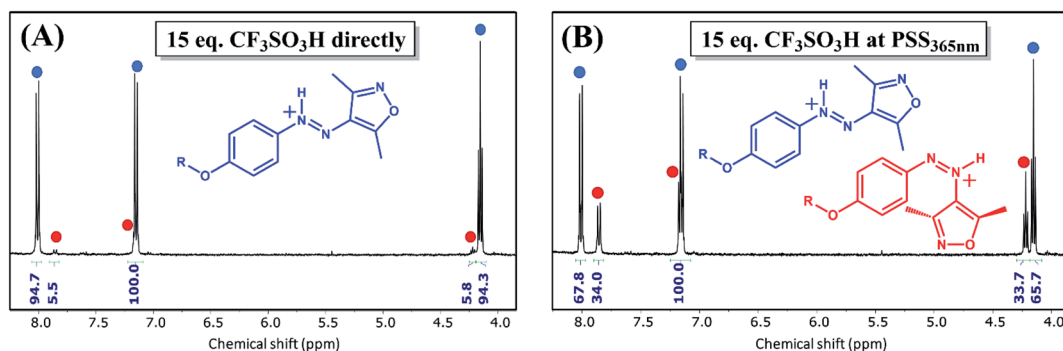


Fig. 4 ¹H NMR spectroscopy of AIZ 3 (3.3 mM in CD_3CN) (A) after immediate addition of 15 eq. of $\text{CF}_3\text{SO}_3\text{H}$ and (B) after addition of 15 eq. of $\text{CF}_3\text{SO}_3\text{H}$ to the PSS_{365nm} (radiant flux 1.2 W). Although the required number of $\text{CF}_3\text{SO}_3\text{H}$ equivalents to form the azonium species is much smaller at ¹H NMR than UV/vis concentration, such stronger interaction at higher concentration ranges has been noted before.³⁷ Note that the delay between addition of $\text{CF}_3\text{SO}_3\text{H}$ and measuring causes the predominantly present *Z*-isomer at the PSS_{365nm} to revert largely to the *E*-azonium. For a full comparison with chemical shifts of the unprotonated forms see Table S1 and Fig. S21†.



indicative of the *E*-and *Z*-azonium observed in UV/vis spectroscopy. Additionally, double protonation would result in blue-shifting of absorptions (Fig. S23†), and no such features are observed upon addition of a large range of acid equivalents (Fig. 5). Instead, an interplay between protonation/deprotonation and reversible *Z/E*-photo- and thermal isomerisation opens up to four distinct regimes of kinetic control in this acidity range. At near-stoichiometric equivalents primarily *E*-AIZ and *Z*-AIZH⁺ are observed (Fig. 5A), differing from catalysis with weak acids as it transiently yields a strongly absorbing *Z*-AIZH⁺ isomer upon UV-irradiation (Fig. 1B). Upon increasing the equivalent CF₃SO₃H to *ca.* 30 eq. the absorption of the red-shifted *Z*-AIZH⁺ reaches a fully photo- (Fig. 5B) and thermoreversible (Fig. S24†) steady state, while a further increase to *ca.* 500 eq. leads to a more significant share of *E*-AIZH⁺ and a limitation of absorption changes primarily to the UV-range (Fig. 5C). At *ca.* 800 eq. of CF₃SO₃H, the absorption is dominated by the azonium species, their large overlap in absorption providing subtle photoreversible changes in the UV- and visible range (Fig. 5D). Thus, we find that although the molecular

components of the abovementioned four photochemical systems are chemically identical, the relative composition provides tuneable control over various absorptions without the need for synthetic alterations.

Acid-catalysed liquid-to-solid phase transitioning

Although observed in few AAPs also,^{51,52} AIZs exhibit a much higher tendency to undergo solid state switching with reversible solid-to-liquid phase transitions, owing to a drastic change in melting point upon the out-of-plane bending in the *Z*-form.⁴³ Fascinatingly, the p*K*_a dependence of the liquefied AIZ is completely orthogonal to that in solution, showing for all three AIZs a swift catalytic reversion in response to CF₃COOH vapours, accompanied by a delayed liquid-to-solid phase transition (Video S1†) that was also observed photochemically.⁴⁴ Analogous to the solution behaviour, CH₃COOH fumes yield no response in phase as well as absorption, while CF₃SO₃H vapours are found to be sufficiently acidic to protonate either isomer, merely resulting in slow *Z/E*-equilibration (Video S2†). Slow dispersion of the CF₃SO₃H in air occurs over time, however,

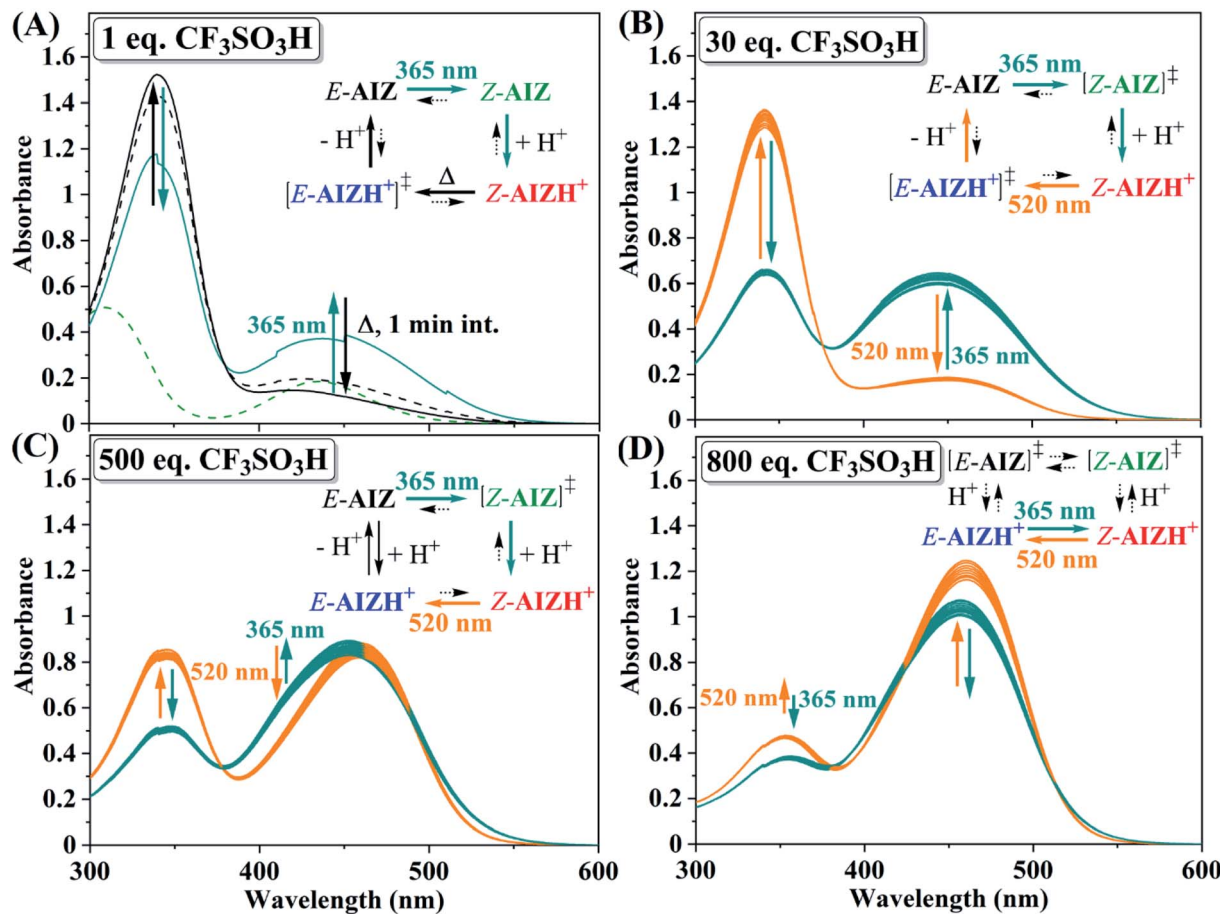


Fig. 5 UV/vis absorption spectra of AIZ 3 (70 μ M in acetonitrile) giving rise to varying photocontrol over azo-speciation at 365 nm (radiant flux 1.2 W, teal arrows) and at 520 nm (radiant flux 87 lm, orange arrows) depending on the amount of CF₃SO₃H present. 20 steps shown for (B)–(D). \ddagger = not clearly observed. Since only 2 species are predominant at 30 and 800 equivalents of CF₃SO₃H, this allows for approximation of the PSSs under these circumstances by comparing the combined absorption at $\lambda = 340$ and 460 nm with those observed in Fig. 1C: (B) PSS_{365nm} \approx 62% *Z*-AIZH⁺ and 38% *E*-AIZ, PSS_{520nm} \approx 20% *Z*-AIZH⁺ and 80% *E*-AIZ; (D) PSS_{365nm} \approx 37% *Z*-AIZH⁺ and 63% *E*-AIZH⁺, PSS_{520nm} \approx 12% *Z*-AIZH⁺, 88% *E*-AIZH⁺.



ultimately leading to resolidification over *ca.* 45 min as stoichiometric quantities of acid still induce catalysed reversion (Fig. 5A). Only trifluoroacetic acid and hydrochloric acid (Video S2†) cause an immediate red-shift in absorption followed up by a blue-shift (Fig. S25†), indicative of transient formation of the metastable red-shifted Z-AIZH⁺ isomer.

Acid-catalysed adhesion

The retention of acid-catalysis in the liquefied Z-isomer followed by the delayed recrystallisation of the E-isomer opens up further opportunities in the application of AIZs as adhesives (Fig. 6A). The delay in nucleation of several seconds after the near-instant acid-catalysis with CF₃COOH vapours provides time to press the surfaces together, while the resulting flux of still liquid E-isomer favourably delays the recrystallisation further and allows for quick readjustments still (Video S3†). Accordingly, the acid-catalysed curing yields glass-to-glass adhesive strengths of $25.8 \pm 4.8 \text{ N cm}^{-2}$ and PMMA-to-PMMA adhesion at $21.9 \pm 2.5 \text{ N cm}^{-2}$ compared to $29.7 \pm 0.1 \text{ N cm}^{-2}$ and $22.9 \pm 1.2 \text{ N cm}^{-2}$, respectively, that were achieved

photochemically.⁴⁴ The larger error margin is due to the adventitious coverage of the acid-catalysis sometimes leaving residual Z-isomer, most notably upon the more rapid acid catalysis with HCl which yields an overall lower glass-to-glass adhesive strength of $15.9 \pm 3.0 \text{ N cm}^{-2}$. The photochemical debonding in the acid-cured adhesives is fully retained, nevertheless, with identically swift response times of photochemical debonding, *i.e.*, *ca.* 0.5 seconds, under near-maximum loads (Fig. 6B, Video S4†). To further probe the influence of acid on adhesive strength, introduction of CF₃COOH during complete thermal liquefaction and resolidification, *i.e.*, adding vapours after heating past the AIZ melting point and immediately conjoining the slides, was found to yield a thermal adhesive strength of $13.9 \pm 1.1 \text{ N cm}^{-2}$. Although the thermal adhesion of 19.6 N cm^{-2} without acid vapours was previously also shown to be lower than in the photochemical pathway,⁴⁴ the adhesive strength is affected by acid vapours more strongly in the thermal pathway. Thus it seems that the efficiency of the catalytic reversion plays a key role in attaining adhesive strengths nearing those obtained photochemically.

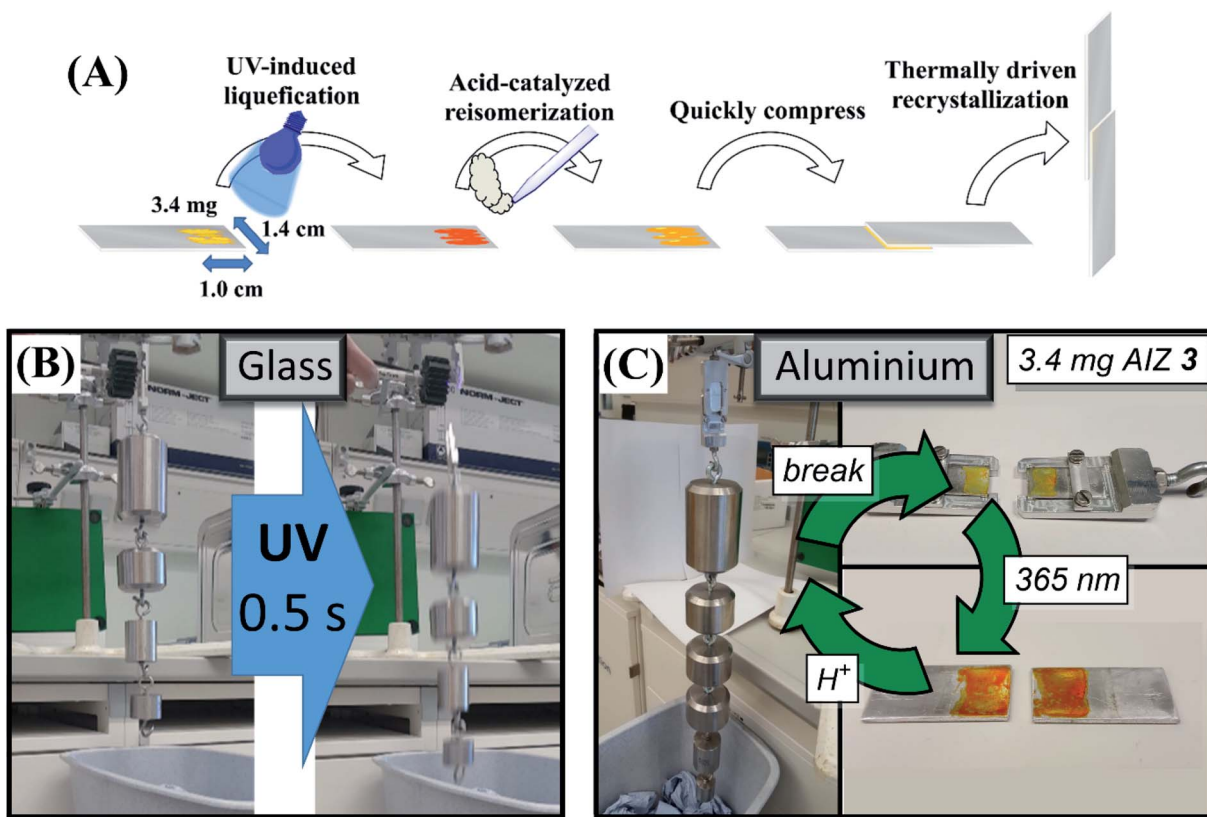


Fig. 6 (A) Strategy for acid-catalysed adhesion independent of the transparency of substrates. Acid-catalysis by CF₃COOH acid vapours causes near-instant Z-to E-reisomerisation of photoliquefied AIZ 3. The several seconds that separate the catalytic recovery of E-AIZ and the advent of nucleation sites for recrystallisation provide time to press slides together before the solidification takes place. (B) Suspension of glass slides conjoined with 2.2 mg of compound 3 at a 1.0 by 1.4 cm overlapping surface area, resulting in an adhesive strength of $25.8 \pm 4.8 \text{ N cm}^{-2}$ (shown here to hold 3.8 kg). The UV-induced debonding⁴⁴ is unaffected by the acid-catalysis over 5 cycles of acid-catalysed bonding and photochemical debonding (Video S4†), fully retaining response times of *ca.* 0.5 s at near-maximum loads. (C) Suspension of aluminium slides conjoined with 3.4 mg AIZ 3 at a 1.0 by 1.4 cm overlapping surface area, capable of holding a load of $35.8 \pm 3.5 \text{ N cm}^{-2}$ (over 15 samples, shown here to hold 5.7 kg). Although the non-transparency rules out photochemical annealing, acid-catalysed bonding can be repeatedly performed at $35.0 \pm 2.8 \text{ N cm}^{-2}$ over 5 cycles of physical debonding.



Interestingly, the lingering presence of trace acid vapours between adhered slides also invokes a self-healing mechanism in the AIZ adhesive (Video S5†), persisting after 10 days still (Video S6†). The debonding is found to be unaffected by the self-healing, as the latter occurs over a longer timescale than the near-instant photoliquefaction of the interfacial thin layer that dictates the adhesive strength. The mechanism of the acid-catalysed reversion thus additionally holds promise for increased resilience against exposure to ambient UV. Furthermore, while the photoreversible nature of the solid-to-liquid phase transition requires the use of at least transparent surface to harden the versatile adhesive at a range of metallic and polymeric substrates,⁴⁴ the acid-catalysed recrystallisation is wavelength independent and opens up to non-transparent surfaces being conjoined also.

In this way, the acid-catalysed pathway further broadens the scope to completely non-transparent substrates with, *e.g.*, a pair of aluminium slides being able to bear $35.8 \pm 3.5 \text{ N cm}^{-2}$ (Fig. 6C, S26†). Though the adhesion of non-transparent surfaces cannot be reversed with light, physical separation of such surfaces (*i.e.*, breaking of the cohesion by force, or heating over the melting point of the adhesive⁴⁴) still yields the chemically unaltered AIZ 3. Accordingly, a pair of aluminium slides was repeatedly glued together to successfully yield an adhesive strength of $35.0 \pm 2.8 \text{ N cm}^{-2}$ over 5 bonding/debonding cycles without increasing the amount of adhesive agent.

Conclusions

Previously, the arylazoisoxazole class of azo switches was shown to possess promising photophysical characteristics in solution as well as the solid-state. The solid-state *E*-to *Z*-AIZ photo-switching accompanied by a reversible solid-to-liquid phase transition made the photoswitch extremely suitable for application in fully photocontrolled reversible adhesion. Here, we demonstrate that the acid-catalysed *Z*-to *E*-AIZ reversion in solution is tuneable by the pK_a of the acid employed. We show that although such acid-catalysis was observed for azobenzenes and azopyrazoles before, the key to this reactivity is that the acid strength is intermediate to the *Z*-and *E*-isomer. Commonly employed acids possess a pK_a lower than only that of *Z*-AIZH⁺, thermodynamically driving the isomerisation over the significantly lower *Z*-AIZH⁺ to *E*-AIZH⁺ energy barrier as confirmed by DFT. Conversely, when an acid is employed that possesses a pK_a that is lower than both the *Z*-and *E*-isomer, such as CF₃SO₃H, it leads to a gradual establishment of the equilibrium between the now thermally stable *E*-AIZH⁺ and *Z*-AIZH⁺, giving rise to several regimes of photoreversible control over a 4-state switching cycle. Most interestingly, the acid catalysis observed in the solution studies applies to the pristine liquefied *Z*-isomer also, providing a chemical trigger for rapid and wavelength-independent hardening of the adhesive with HCl or CF₃COOH vapours. Moreover, although the *E*-isomer has no strong enough interaction with CF₃COOH vapours, traces of the acid remain after the acid-catalysed adhesion and provide a continual self-healing mechanism without distorting the adhesive properties. The non-destructive chemical hardening of

the adhesive and the fundamental understanding thereof opens up further opportunities in the use of the azoisoxazole class of photoswitches, not least the successful adhesion of non-transparent substrates. Furthermore, the adhesive shows excellent reusability after physical breaking of cohesion between non-transparent substrates, retaining its “green” application. Ultimately, the expanded understanding of the acid-reactivity in azoisoxazoles and the demonstrated advance in application have broad implications for the pH-tunable functionality of the myriad of existing azobenzenes and azoheterocycles.

Data availability

The datasets supporting this article have been uploaded as part of the ESI.

Author contributions

LK conceptualised the work and refined it with the help of JS, NBA and BJR. JS performed the synthesis and characterisation of compounds 1–4. LK performed the primary investigation, NBA supported on the solution studies. MB performed the theoretical investigation under supervision of NLD. BJR provided supervision and reviewed and edited the draft, LK wrote the manuscript.

Conflicts of interest

There are no conflicts to declare.

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