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Introduction

Photoswitching is an intramolecular transformation between two states triggered by light absorption.^{1–5} Based on the photochrome structure, photoswitches are grouped into prototypical families such as azobenzenes,^{5,6} stilbenes,⁷ spiropyrans,⁴ indigos,⁸ diarylethenes,⁹ fulgides,¹⁰ and (acyl)hydrazones.^{2,11} Photoswitching most commonly occurs *via* isomerization of a double bond or electrocyclisation.¹ The character of this transformation translates into property differences between the two switched forms and hence into the application potential. Configurational switching has been used for catalytic site shielding,¹² allosteric activation,¹³ photocomplexation^{14,15} and photopharmacology.^{16,17} Electrocyclizations change the

Closed-to-open-shell ground state photoswitching of thienyl-based acylhydrazones†

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Photoswitches convert a photon to molecular information through a reversible change in bonding or configuration. Applications of photoswitches are determined by their different properties in the two states, spanning from catalysis, to allosteric activation, to photopharmacology, up to materials and devices such as memories that can store optical information in chemical bonds. For the latter, operation in the solid state and different means to read/write/erase the information must be possible. These features, however, are challenging to achieve due to close molecular packing and intermolecular interactions. Here, we present acylhydrazones that photoswitch even in the solid state and adopt an open-shell triplet ground state electronic configuration in the photoswitched form, which is very longlived. The optically stored information can thus be read magnetically and completely erased by applying an electrochemical potential. In addition, ab initio calculations identified the closed-to-open-shell pathway and explained the singlet-triplet crossing. Photoswitching of the closed/open-shell ground state electronic structure is a fundamentally new feature, which can bring new opportunities and applications. We demonstrate it in an example of the open-shell state of the photoswitch used as an initiator of radical benzylic bromination and radical debromination of an aromatic ring. However, the presented phenomenon is robust and insensitive to conditions, thus showing potential in a broad range from heterogeneous catalysis to spintronics.

> conjugated backbone and finds applications in (organo)catalysis,^{18,19} smart materials,^{4,20,21} and optical memories.^{3,9,10} For memories, however, the read-out of the information should occur by non-optical means so that secondary light does not alter the stored information. Electrical²² or magnetic²³ reading could be an option if the magnetic or electronic properties change significantly on photoswitching, *i.e.* if the two forms have different electronic configurations, such as closed- and open-shell. Moreover, a radical form of open-shell species is highly interesting for the initiation and catalysis of reactions.

> Acylhydrazones are known to perform reversible photoisomerisation of the C=N bond between the more thermodynamically stable *E*-form and the *Z*-form.^{2,24-26} The reverse isomerization ($Z \rightarrow E$ backswitching) can occur both thermally (T-type photoswitches) or exclusively by excitation at different wavelengths (P-type), depending on the isomerization energy barrier and spectral separation of the two forms.

> P-type switching is particularly interesting for optical information storage^{3,9,10} and catalysis because the properties associated with the Z-form do not revert at a significant rate at a given temperature.^{11,21,27} Further properties are required for application in practical devices: (1) the information must be written reliably, *i.e.* with the highest possible quantum yield;³

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(2) the information reading must not alter the stored information, that is, the quantum yield of backswitching should be much lower or not occur during the reading process;²⁸ and (3) the photoswitching properties must be achieved in the solid state.^{29–31} For catalysis, additional features such as stability under the reaction conditions and solubility are required. Achieving solid-state photoswitching is a paramount challenge due to efficient intermolecular quenching of excited states and the high steric hindrance imposed by molecular close packing.

Here, we present a series of acylhydrazones with the thiophene moiety in the azomethine part of the molecules to (1) form long-lived open-shell radicals in their photoswitched form, (2) erase the stored information by an electric impulse, and (3) operate in the solid state. This alone demonstrates the potential of the series of acylhydrazones for application in solid-state devices. We further show their application in catalysis.

Results and discussion

Acylhydrazones bearing the 2-thienyl group on the azomethine side have been reported² to exhibit a large bathochromic shift of absorbance in the photoswitched form, so that both forms can be irradiated specifically without the contribution of the other. The large bathochromic shift is particularly intriguing because both *E*- and *Z*-forms are anticipated to be rather planar and thus no dramatic change in conjugation is expected. Therefore, the anticipated planarity of the *Z*-state is in dispute and hereafter we address this state as the 'photoswitched form' to discuss its geometry below. Synthesis of acylhydrazones is a trivial procedure (Fig. 1), providing the corresponding products in high yields. The starting compounds are available commercially at low cost or can be readily synthesized by conventional methods. The ease of procurement allows straightforward scale-up.

The series of prepared derivatives (Fig. 1) showed similar photophysical properties, *i.e.* a large bathochromic shift of



Fig. 2 UV-vis spectra of AH5 recorded during irradiation at the 340 nm absorbance maximum. Similar results are observed for all 2-thienyl based acylhydrazones. Excellent separation of new absorbance features associated with the photoswitched form was confirmed in the whole series (see the ESI† for experimental traces of other derivatives).

absorbance on photoswitching, negligible thermal back conversion in the solid state, and slow back-isomerisation by visible light for all 2-thienyl-based acylhydrazones in acetonitrile solution (**AH1**, **AH3-9**; see also the ESI†). The absorbance maxima in both *E*- and photoswitched forms shift bathochromically with extended conjugation from UV to the blue visible range (400–450 nm) in the case of thienothiophene derivatives (Fig. 2). Furthermore, excellent switching repeatability was observed in the case of **AH1** and **AH3-9** (see the Photoswitching section, ESI†).

AH2 was an exception to the series. It did not show any new absorbance features in the UV-vis spectra upon irradiation and therefore the *Z*-form cannot be specifically addressed photochemically (Fig. S6, ESI†). **AH7** and **AH8** could also be predicted to be different due to potential two-stage photoswitching. However, UV-vis spectra recorded during irradiation show isosbestic points. This suggests that only one of the acylhydrazone



Fig. 1 Scheme of acylhydrazone synthesis and the list of prepared photoswitches with their acronyms used in this study.

moieties can be isomerized even at prolonged irradiation times as the UV-Vis spectra do not evolve after 5 minutes of irradiation (Fig. S25 and S29, ESI[†]). This is supported also by the ¹H NMR spectrum, in which the loss of symmetry leads to the emergence of a new set of signals (Fig. S27, ESI[†]). The ¹H NMR spectra remained the same even at a prolonged irradiation time (12 h). We have determined the quantum yield for the $E \rightarrow$ photoswitched-form isomerization of all AHs (*e.g.* the quantum yield of **AH1** is 0.39; see Section S4, ESI[†]), which is in line with other reported acylhydrazone photoswitches.²

The photoswitching properties obtained in acetonitrile solutions motivated us to study our set of acylhydrazones in the solid state. Thin films prepared by drop-casting acylhydrazone AH1-AH9 solutions indeed showed reversible photoswitching (see the ESI[†] for solid-state photoswitching). The anticipated severe steric demands of switching in the solid state were shown to be less significant by the XRD single crystal structure (Section S12, ESI[†]). Crystal packing showed void volumes around the phenyl group of the hydrazide part. Importantly, the C=N $E \rightarrow$ photoswitched-form transformation can occur through flipping of the carbonyl moiety, accompanied by rotation around the carbonyl-phenyl σ -bond without actual rotation of the benzene ring. In this scenario, most of the steric obstacles are obviated and the free space around the phenyl group provides the necessary degree of freedom to facilitate the flipping of the rather small = N-NH-(CO)- group. However, despite our efforts, we did not achieve photoswitching of the single crystal nor were we able to obtain a monocrystal of the photoswitched form.

Due to scattering in solid films of the photoswitches causing significant spectral broadening, we have prepared 'solid solutions' of **AH1–9** in a poly(methyl methacrylate) (PMMA) matrix and repeated photoswitching studies (see Section S6, ESI,† for photoswitching of AHs in the PMMA matrix). In all cases, the recorded spectra followed the same trends as those of acetonitrile solutions, *i.e.* a new bathochromic absorbance of

photoswitched isomers, a long-lived photoswitched state, and slow back-isomerization by visible light. These experiments have two important implications. First, steric hindrance and intermolecular quenching of excited states in the crystal packing or polymer matrix do not prohibit switching of these acylhydrazones. Second, the bathochromically shifted absorbance feature of the photoswitched form is of the monomolecular origin because excimer formation is difficult to imagine in the PMMA matrix.

We used cyclic voltammetry (CV) and in situ UV-vis spectroelectrochemistry to clarify the particular behavior of 2-thienylbased acylhydrazones. CV showed up to three irreversible oxidation waves in the range of 1.6 to 2.0 V for all acylhydrazones (Section S7, ESI⁺), and double acylhydrazones AH7 and AH8 also exhibited a reversible reduction peak at around -0.4 V. For spectroelectrochemistry, we irradiated the acylhydrazone solution in acetonitrile with light (wavelength adjusted to the most bathochromic absorbance maximum) to isomerize to the photoswitched form. After reaching the photostationary state (PSS), we performed cyclic voltammetry and monitored the UV-vis spectra. When a CV cycle of 0 V \rightarrow 2.5 V \rightarrow -2.0 V \rightarrow 0 V was applied, a dramatic and rapid change observed in the UV-Vis spectrum indicated complete reversal of the photoswitched form to the E-form (Fig. 3, kinetics in the ESI[†]). This behavior is common to all 2-thienyl-based acylhydrazones in the studied series. Switching under cyclic voltammetry is interesting, as it shows that the optical information stored in the configuration of the molecule can be efficiently erased by applying an electrochemical stimulus. In essence, AHs act as "electroswitches". We attribute the switching to the radical cation propagation in the medium, which has been described on azobenzenes.32

The body of data obtained from spectroscopy and electrochemistry of the photoswitched form led us to suspect an openshell electron configuration. We have employed electron paramagnetic resonance (EPR) studies in solution in the presence of

Fig. 3 UV-Vis spectra of AH5 photoisomerization (black to blue) and the spectrum after the CV cycle (red). The CV cycle led to a dramatic and instantaneous change in the UV-Vis spectrum, indicating a complete reversal to the *E*-form. The 1 H NMR spectrum shows that only the photoswitched form of AH5 was transformed back to aldehyde after CV.

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Fig. 4 EPR spectrum (X-band) of the trapped radical species between irradiated AH5 and the spin trap agent in acetonitrile. Concentrations were set to c(AH5) = 10 mM and c(N-t-butyl- α -phenylnitrone) = 45 mM. The EPR spectrum was centered at $g_{iso} = 2.0065$. The hyperfine splitting structure comes from $a (1 \times {}^{14}\text{N}) = 1.35 \text{ mT}$ and $a (1 \times {}^{14}\text{H}) = 0.22 \text{ mT}$.

a spin trap. To investigate this, **AH5** was converted from *E*- to the photoswitched form in the presence of a spin trap (*N*-*t*-butyl- α -phenylnitrone, by irradiating with 340 nm light). In the resulting solution, an EPR spectrum gave a clear signature of a trapped radical species (Fig. 4). The solution without the trap agent did not show any detectable signal. Therefore, the magnetic properties of the compound can be "read" by

an independent method (EPR) from the one that caused photoswitching.

To shed light on the electronic structure and transitions, calculations of the ground and excited states were performed. All vertical excitation energies were calculated using the Complete Active Space Second Order Perturbation Theory (CASPT2) method, with the zero-order wavefunction obtained with state-averaged complete active space (SA-CASSCF) on AH1 and AH2. These two systems are the smallest in the investigated series and thus accessible at such a high level of calculations. Both AH1 and AH2 were investigated to compare the molecules that exhibit photoswitching associated with the bathochromic absorbance shift (AH1) with those that do not (AH2).

The **AH1** ground state optimization indicated a mix of two ground state geometries close in energy, differing only in thiophene ring conformation (*s-cis* and *s-trans*, Fig. S75, ESI†). Calculation of vertical transitions to S_{1-3} produced results consistent with the experimentally measured UV-Vis spectra (Tables S1, S2 and Fig. S1, ESI;† summarized in Fig. 5). Relaxation of these excited states follows three pathways leading to local conformational minima (Table S3, ESI,† "radiative geometry 1", "radiative geometry 2" and "switching geometry" in Fig. 5). The radiative geometries have a very similar torsion angle along the C=N bond to the ground state structure (Tables S3 and S7, ESI†). Their relaxation by photon emission leads to a fluorescence spectrum that agrees very well with the experiment (Fig. 5).

Fig. 5 The proposed scheme for the photoswitching and radiative decay of AH1 upon excitation from the lowest energy ground state geometry to the S1–S3 states. Optimization of these states pointed at three geometries: "radiative geometry 1", "radiative geometry 2" and "switching geometry" shown in the bottom center. In the "switching geometry", SOC allowed S0 \rightarrow T1 crossing, followed by further relaxation to T1 minima. Various energy differences are highlighted with their corresponding wavelengths and compared to experimental UV-vis spectra (top left) and fluorescence spectra (bottom left). T1 \rightarrow T2 excitation represents a pathway for the back photoswitching. The molecular orbitals with occupancy close to 1 of the open-shell electronic structure in S1 and T1 are depicted in the bottom right.

The "switching geometry" of **AH1** has some unique properties. Along the S₁ relaxation pathway to the "switching geometry", the torsion angle along the C=N bond changes from almost parallel to basically perpendicular (Table S3, ESI†). In this geometry, the S₀ state most closely resembles the S₁ state in the ground state geometry with both having open-shell character corresponding to an imine $\pi \rightarrow \pi^*$ transition.

It is hence assigned that the ground state of AH1 undergoes vertical excitation to the S1 state in the "ground state geometry", which relaxes to the S₀ state in the "switching geometry" through a conical intersection. It is from here that conversion to the triplet state most likely occurs due to (1) the high $S_0 \rightarrow T_1$ spin-orbit coupling (SOC) of 22 cm⁻¹ calculated in this geometry (Table S4, ESI^{\dagger}), and (2) the lower energy of T₁ than that of S_0 . This T_1 state in the switching geometry relaxes further by 0.46 eV or 0.53 eV to two geometrically almost identical T₁ minima. Calculation of $T_1 \rightarrow T_2$ vertical excitations from these geometrical minima suggested peaks at 3.04 eV and 3.33 eV (408 and 372 nm, respectively), which is in excellent agreement with the observed bathochromically shifted absorbance of AH1 in the photoswitched state. Calculations also revealed a shallow conformational energy surface around these T₁ geometries (Section S9.4, ESI⁺), which supports the observed long lifetime of these triplet species in both solid and solution states. Also, due to this conformational flexibility, describing the photoswitching of 2-thienyl-based acylhydrazones as $E \leftrightarrow Z$ isomerization is an oversimplification, although generally used also for azobenzenes.³³ Hence, we assign the AH1 photoswitching to conversion of the closed-shell E-isomer into the triplet ground state photoswitched form adopting a perpendicular orientation of the azomethine and hydrazide parts of the photoswitch. The process of conversion is summarized in Fig. 5.

Calculations of AH2 show important differences. Relaxation from the vertically excited S₁ state also showed pathways to three possible geometries. However, the C=N torsion angle does not approach perpendicularity in any of them. In contrast to AH1, for all S1 minima geometries identified for AH2 (Table S5, ESI[†]), the lowest electronic states reveal a closedshell character. Furthermore, the S1 minima of AH2 is the most similar in geometry to that in AH1, which does not have the inversion of the closed shell state and the open shell (imine $\pi \rightarrow \pi^*$) state. Finally, although the calculated spin-orbit coupling elements at these geometries are relatively large, suggesting potential intersystem crossing, the energy ratios favor decay to the system with a closed-shell structure rather than a structure with radical character. Thus, in the case of AH2, the photoswitching occurs between closed-shell species and can be described as a plain $E \leftrightarrow Z$ process. The observed decrease of absorbance during photoswitching is consistent with the calculated lower oscillator strength of the Z-isomer.

Localization of singly occupied orbitals in **AH1** at both T₁ minimum and S₀ in the switching geometry shown in Fig. 5 indicated separation of radicals on the two constituents, thiophene ring and the hydrazide. These constituents are very close to each other in the molecule. Likely due to the proximity and delocalization of the two spin sites (radical centers), we could not detect any $\Delta M_s = \pm 2$ EPR (double quantum) signal characteristic of the triplet electronic configuration. This also explains why we were able to record clean NMR spectra of the photoswitched forms; although radicals are generally considered detrimental to NMR spectroscopy characterization, triplet states can provide sharp and easy-to-interpret spectra depending on the spin coupling. Furthermore, NMR experiments with an internal standard (*t*-BuOH) showed that all species

Fig. 6 (a) Catalysis of radical halogenation to the benzyl position by AH5 in a photoswitched form. (b) The new system featuring both an AH double bond and a DAE ring, which can be closed and opened upon irradiation and can catalyze radical dehalogenation in its photoswitched form.

originating from photoswitching, *i.e.* the *E*- and the photoswitched form, are observable in the NMR spectrum (Fig. S4, ESI[†]).

Therefore, we made an effort to separate the two radical centers and designed a new system - DAE-2AH1 (Fig. 6), which combines two AH1 motifs with a diarylethene (DAE) photoswitch (bis-thienyl in our case). DAE undergoes photochemical electrocyclization, which brings the two decoupled aromatic rings in the open state in conjugation in the cyclized form.^{9,20} We substituted the two unoccupied α -positions of the two thiophene rings with AH1 motifs by grafting aldehyde functions followed by condensation with benzhydrazide. Upon irradiation with 340 nm light, we observed a gradual development of a strong purple color characteristic of the closed form of DAE. Using NMR and UV-vis, we also confirmed that both AH1-sites were converted to the photoswitched state as well. In this photoswitched form, the two hydrazide parts of DAE-2AH1 are isolated by the torsion angle around the C=N bond, but the two thiophene moieties are conjugated. This allows the thiophene singly occupied states to recombine and therefore the two spin sites on hydrazone moieties are separated from each other in space.

The triplet character of the photoswitched form of AH5 acts as a diradical, which inspired us to use it as an initiator in a model radical reaction - radical bromination of toluene in the benzylic position by N-bromosuccinimide (NBS) (Section S11, ESI[†]). We mixed AH5 either in the *E*-form or in the photoswitched form with toluene and NBS in acetonitrile and monitored the conversion of toluene to benzyl bromide by HPLC for three hours. The reaction was conducted in the dark to avoid secondary irradiation effects. Subsequently, the solvent was evaporated, and the product formation was confirmed by NMR. Reaction without any form of AH5 served as a negative control. While no benzyl bromide was detected in the negative control after three hours, the run with as-synthesized AH5 showed 11% conversion, and the run with AH5 pre-irradiated to the PSS reached 60% conversion over the same period. This clearly demonstrates that the triplet state lasts sufficiently long even under the conditions of radical bromination by NBS and acts as an efficient initiator of the reaction.

Based on the above-mentioned successful application of **AH5**, we decided to demonstrate also the power of **DAE-2AH1** to drive radical reactions. Radical dehalogenation of 2-bromoanisole was chosen as a model reaction. We mixed **DAE-2AH1** in the photoswitched form with 2-bromoanisole and tributylstannane in DMF and monitored the conversion of 2-bromoanisole to anisole by HPLC. The reaction was also conducted in the dark to avoid secondary irradiation effects. The presence of the product was confirmed by comparison with the standards by HPLC and also by ¹H NMR. We also used ¹H NMR to prove that the reaction without **DAE-2AH1** does not yield the product.

Conclusions

We synthesized a series of thienyl-based acylhydrazones in high yields using the reaction of selected hydrazides and aldehydes (AH1-9). We observed a similar photoswitching of all 2-thienyl derivatives. Furthermore, we observed strikingly different photoswitching in the 3-thienyl derivative. Photoswitching is repeatable and the photoswitched form is stable according to the long half-life, which is above 6 hours. Solid-state photoswitching was observed in pure films and solid solutions in a PMMA matrix. The optically stored information is completely erased electrochemically at reasonable potentials. We attributed the specific spectral features of photoswitched isomers of 2-thienyl acylhydrazones to the open-shell triplet configuration by EPR and ab initio calculations. Further calculations provided excellent agreement of experimental spectra with calculated predictions for triplet states. Thorough theoretical investigation of excited states and their geometry optimization by multireference methods allowed us to describe the isomerization pathway. Finally, we demonstrated the application potential of the presented acylhydrazone systems in catalysis as radical initiators. The closed-to-open-shell reversible photoswitching is an unprecedented phenomenon allowing manipulation with spin multiplicity between two long-lived states. This opens up unique opportunities in many fields such as catalysis, information storage, spintronics, and materials science.

Author contributions

P. K. designed the study. M. Š., A. G. and O. M. synthesized the compounds. M. Š., V. G., H. R. and O. M. performed spectroscopic and electrochemical experiments. M. Š. and V. G. determined the quantum yields. A. L. and D. N. designed and performed the *ab initio* calculation study. J. T. performed, analyzed and interpreted the EPR data. V. E. measured, analyzed and interpreted the XRD data. M. Š. performed the catalytic study. M. Š., H. R. and P. K. consolidated all data, discussed the results and cowrote the manuscript. All authors contributed to the manuscript editing.

Data availability

All the data is available in the repository under DOI: 10.5281/ zenodo.14681032.

Conflicts of interest

The authors declare no competing interests.

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References

- 1 R. Göstl, A. Senf and S. Hecht, *Chem. Soc. Rev.*, 2014, 43, 1982–1996.
- 2 D. J. van Dijken, P. Kovaříček, S. P. Ihrig and S. Hecht, *J. Am. Chem. Soc.*, 2015, **137**, 14982–14991.
- 3 M. Irie, Chem. Rev., 2000, 100, 1683-1684.
- 4 R. Klajn, Chem. Soc. Rev., 2013, 43, 148-184.
- 5 G. S. Hartley, Nature, 1937, 140, 281.
- 6 E. Merino, Chem. Soc. Rev., 2011, 40, 3835-3853.
- 7 A. Kanaya, Y. Takashima and A. Harada, *J. Org. Chem.*, 2011, **76**, 492–499.
- 8 C.-Y. Dennis Huang and S. Hecht, *Chem. Eur. J.*, 2023, **29**, e202300981.
- 9 M. Irie, T. Fukaminato, K. Matsuda and S. Kobatake, *Chem. Rev.*, 2014, **114**, 12174–12277.
- 10 Y. Yokoyama, Chem. Rev., 2000, 100, 1717-1740.
- 11 G. Men and J.-M. Lehn, Chem. Sci., 2019, 10, 90-98.
- 12 S. Yang, D. Larsen, M. Pellegrini, S. Meier, D. F. Mierke, S. R. Beeren and I. Aprahamian, *Chem*, 2021, 7, 2190–2200.
- 13 D. Hager and D. W. C. MacMillan, J. Am. Chem. Soc., 2014, 136, 16986–16989.
- 14 G. Vantomme and J.-M. Lehn, Angew. Chem., Int. Ed., 2013, 52, 3940–3943.
- 15 H. Fu, S. Pramanik and I. Aprahamian, J. Am. Chem. Soc., 2023, 145, 19554–19560.
- 16 W. A. Velema, W. Szymanski and B. L. Feringa, J. Am. Chem. Soc., 2014, 136, 2178–2191.
- 17 P. Kobauri, F. J. Dekker, W. Szymanski and B. L. Feringa, Angew. Chem., Int. Ed., 2023, 62, e202300681.
- 18 R. Dorel and B. L. Feringa, *Chem. Commun.*, 2019, 55, 6477–6486.

- F. Eisenreich, M. Kathan, A. Dallmann, S. P. Ihrig, T. Schwaar, B. M. Schmidt and S. Hecht, *Nat. Catal.*, 2018, 1, 516–522.
- 20 M. Kathan, P. Kovaříček, C. Jurissek, A. Senf, A. Dallmann,
 A. F. Thünemann and S. Hecht, *Angew. Chem., Int. Ed.*, 2016, 55, 13882–13886.
- 21 M. Kathan, C. Jurissek, P. Kovaříček and S. Hecht, J. Polym. Sci., Part A: Polym. Chem., 2019, 57, 2378–2382.
- 22 I. Hnid, D. Frath, F. Lafolet, X. Sun and J.-C. Lacroix, *J. Am. Chem. Soc.*, 2020, **142**, 7732–7736.
- N. E. Beyrouti, F. Houard, M. Cordier, E. Trzop, S. Rigaut,
 B. L. Guennic, K. Bernot and L. Norel, *Chem. Commun.*, 2023, 59, 5265–5268.
- 24 M. N. Chaur, D. Collado and J.-M. Lehn, *Chem. Eur. J.*, 2011, **17**, 248–258.
- 25 G. Vantomme, S. Jiang and J.-M. Lehn, J. Am. Chem. Soc., 2014, **136**, 9509–9518.
- 26 X. Su and I. Aprahamian, Chem. Soc. Rev., 2014, 43, 1963–1981.
- 27 J. Holub, G. Vantomme and J.-M. Lehn, J. Am. Chem. Soc., 2016, 138, 11783–11791.
- 28 M. Alemani, M. V. Peters, S. Hecht, K.-H. Rieder, F. Moresco and L. Grill, *J. Am. Chem. Soc.*, 2006, **128**, 14446–14447.
- 29 R. Falkenburg, M. J. Notheis, G. Schnakenburg and L. K. S. von Krbek, Org. Biomol. Chem., 2023, 21, 4993–4998.
- 30 N. Eleya, S. Ghosh, E. Lork and A. Staubitz, J. Mater. Chem. C, 2021, 9, 82–87.
- 31 C. Knie, M. Utecht, F. Zhao, H. Kulla, S. Kovalenko, A. M. Brouwer, P. Saalfrank, S. Hecht and D. Bléger, *Chem. – Eur. J.*, 2014, **20**, 16492–16501.
- 32 K. Munkerup, D. Romanov, T. Bohinski, A. B. Stephansen,
 R. J. Levis and T. I. Sølling, *J. Phys. Chem. A*, 2017, 121, 8642–8651.
- A. Mostad, Chr Rømming, Chr Rømming, S. Hammarström,
 R. J. J. C. Lousberg and U. Weiss, *Acta Chem. Scand.*, 1971,
 25, 3561–3568.