A rechargeable molecular solar thermal system below 0 °C†

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An optimal temperature is crucial for a broad range of applications, from chemical transformations, electronics, and human comfort, to energy production and our whole planet. Photochemical molecular thermal energy storage systems coupled with phase change behavior (MOST-PCMs) offer unique opportunities to capture energy and regulate temperature. Here, we demonstrate how a series of visible-light-responsive azopyrazoles couple MOST and PCMs to provide energy capture and release below 0 °C. The system is charged by blue light at −1 °C, and discharges energy in the form of heat under green light irradiation. High energy density (0.25 MJ kg\(^{-1}\)) is realized through co-harvesting visible-light energy and thermal energy from the environment through phase transitions. Coatings on glass with photo-controlled transparency are prepared as a demonstration of thermal regulation. The temperature difference between the coatings and the ice cold surroundings is up to 22.7 °C during the discharging process. This study illustrates molecular design principles that pave the way for MOST-PCMs that can store natural sunlight energy and ambient heat over a wide temperature range.

Introduction

Thermal management is crucial in our modern society, regardless of whether we are considering chemical transformations, electronics, human comfort, energy production or our whole planet. Thermal management materials based on specific heat capacity or phase change are seeing increased use in applications such as electronics, and domestic and industrial heat management.\(^1\) Phase change materials (PCMs) are a broad class of materials whose latent heat during a phase transition from solid-to-liquid can be used for energy storage applications. Latent heat storage offers a significant advantage if the application involves temperature cycles close to the melting point since in those cases, the corresponding storage density of sensible thermal storage is small. In building applications, phase change materials made from paraffins, salt hydrates, fatty acids or ice can be used as central heat sinks\(^6\) and also in floors, windows or walls.\(^7\) Common to all “traditional” thermal energy storage materials is that they operate via heat transfer, both in energy input and energy output.\(^8\) This leads to design challenges and scaling factors that restrict practical performance and implementation.

Molecular solar thermal (MOST) systems have been recognized as a promising avenue to harvest and store thermal energy.\(^9\)–\(^13\) In the charging process, a stable isomer of a photochromic molecule absorbs photon energy and is converted into a high-energy metastable isomer, thereby storing solar energy in chemical bonds. The MOST system is discharged when the metastable isomer switches back to the stable isomer by external stimuli, with the release of stored energy in the form of heat. While the MOST system shares some properties with PCMs, the process of energy storage and release in the MOST system is controlled by photons and molecular thermodynamics,\(^14\)–\(^17\) whereas in PCMs it is controlled by heat transfer. Recently, combining the functions of MOST and PCMs into a single component material (MOST-PCM) has been utilized to add storage capacity to the MOST system since the charging of the system is not only happening via solar irradiation but also by taking energy directly from the environment.\(^18\)–\(^21\) This dual input leads to an increased energy density by almost 100%.\(^19\)
Another attractive feature is added to PCMs; since the solidification of the cis liquid is not happening spontaneously, the phase change is locked by the photochemical system. This feature dramatically extends the functionality of the MOST-PCM combination since the phase change is controlled by external stimuli and no insulation is needed to hold the latent heat.

However, a severe limitation of MOST-PCMs based on azo-molecules studied until now is their inability to be charged and discharged in the solid state in cold environments, especially below 0 °C, because of the high melting point ($T_m$) of cis-isomers. This is a critical condition since many applications such as thermo regulated fabrics or functional coatings will need to be able to function at that temperature. Generally, the trans–cis photoisomerization ofazo molecules requires a large free volume and can only occur in the surface layers of trans-crystals, thus preventing the charging process in the neat solid state. If the ambient temperature exceeds the cis-isomer $T_m$, the generated cis-isomer melts into a liquid and exposes new trans-crystal surfaces, and finally the trans-crystals are entirely transformed into cis-liquids. But most reported $T_m$ values of cis-isomers are in the range of 20–200 °C, which means that their photoisomerization from trans-crystals to cis-liquid cannot occur at low ambient temperatures. On the other hand, although some cis-isomers can maintain liquid states below 0 °C due to their supercooling behavior to achieve discharging at low temperature, the charging process is still at room temperature (27 °C), limiting the versatility of the system.

Another challenge for the MOST-PCM is that the charging process generally needs UV light irradiation, since UV light causes damage to materials and the human body, and comprises a small fraction (4.5%) of the total solar spectrum, resulting in the low utilization efficiency of solar energy. To date, only one study has reported the utilization of ortho-functionalized azobenzene derivatives to store both visible light energy and room temperature ambient heat. However, this system could not be charged below 0 °C, and the energy density was in the 0.07–0.15 MJ kg$^{-1}$ range. Ortho-Substitution can increase the energy of trans-isomers or decrease the energy of cis-isomers, so that the $\Delta H_{m}$ of this type of azo molecule decreases to only 6–25 kJ mol$^{-1}$ (0.01–0.05 MJ kg$^{-1}$). Therefore, a reversibly charging/discharging and visible-light-energy storage MOST-PCM working at low-temperature remains to be explored.

Here, we report new arylazopyrazoles as MOST-PCMs, which are rechargeable below 0 °C by visible light, as illustrated in Fig. 1. In addition, by co-harvesting the visible light energy and low-temperature ambient heat, an energy density of 0.25 MJ kg$^{-1}$ is achieved, which is an increase of 67% over previous comparable systems. Furthermore, the cis-isomer has a half-life of 22 days at 0 °C, demonstrating its stable energy storage capacity. The combination of high energy density, storage time, and the fact that the system can be charged at low temperatures provides the opportunity to explore the function of the material in a new type of optically regulated MOST-PCM window as a proof-of-concept study, designed to illustrate the function of the MOST-PCM system in a coating. Glass coated with arylazopyrazoles is prepared as a miniature energy storage window. One point to highlight is that the novel windows can be charged and discharged at −1 °C by 400 nm blue light and 532 nm green light, respectively. During the discharging process, the surface temperature of the window can reach from −1 °C up to 21.7 °C (a temperature increase of 22.7 °C), corresponding to a thermal power output of 256.2 W m$^{-2}$ during a continuous period of 60 s. Charging and discharging energy at low temperatures has potential implications for functional clothing, advanced sunglasses, deicing and home heating under ice-cold conditions, thereby increasing thermal comfort and reducing the energy consumption of conventional heating.

**Results and discussion**

**Synthesis and UV-Vis absorption spectra of bidirectional visible-light-driven photoswitches**

The fundamental principle to realizing low-temperature working and visible-light energy storing MOST-PCM systems is...
to design photochromic molecules that drive phase transition with visible light at low temperature. The introduction of a 4-thiomethyl group and changing the bridging positions between the azo group and pyrazole ring was expected to achieve bidirectional visible light switching through extending the π-conjugation. Accordingly, three 4-methylthioarylazopyrazoles (S3, S4, and S5) were designed and their synthetic routes are shown in Scheme 1.

In previous studies,23 arylazopyrazoles were usually prepared by the Mills reaction of nitrosobenzene analogs and aminopyrazoles, but in this method it was difficult to synthesize arylazopyrazoles with electron-donating groups because the electron-donating groups would destabilize nitrosobenzene analogs.24 Hence, we first prepared 3(5)-nitroso-1H-pyrazole, which was then coupled with aniline analogs to give S3(5)-H. S3 and S5 were subsequently produced in one pot by N-methylation at two selectable positions, benefiting from the proton transfer and tautomerism of 1H-pyrazole. S4 was formed via diazo-coupling and cyclization reactions.

Their photoisomerization behaviors were studied using UV-Vis absorption spectra in an acetonitrile solution. As shown in Fig. 2b–d, all trans-isomers exhibited single and intense absorption bands in the 350–400 nm region ($\lambda_{\text{max}} = 25–32 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$) due to $\pi-\pi^*$ transition. Compared to the reported 4-methoxyarylazopyrazole (O4, 342 nm),35 the $\pi-\pi^*$ $\lambda_{\text{max}}$ of S3 and S4 were red-shifted to about 360 nm, and further to 385 nm for S5 (Table S1†). The trans/cis relative absorption of S3, S4 and S5 at 400 nm was strong (Table S1†), and hence it was possible to realize trans-cis isomerization using visible light. The photostationary states (PSSs) at different wavelengths (from 365 to 532 nm) were studied (Fig. S1†), and the isomeric compositions are presented in Table S2.† As a result, 400 nm blue light induced a near-quantitative yield (>95%) of trans-cis isomerization for S5, while only ~85% for S3 and S4. The higher trans-cis photoconversion of S5 was attributed to its $\pi-\pi^*$ $\lambda_{\text{max}}$ closer to 400 nm and stronger trans/cis relative absorption at $\lambda = 400$ nm. Exciting the tail of n–p* bands of three cis-isomers using green light (532 nm) resulted in cis-trans isomerization. The

Scheme 1 Synthesis routes of 4-methylthioarylazopyrazoles. Reagent and conditions. (a) AcOH, room temperature, 58%. (b) CH3I, CS2CO3, DMF, room temperature, 89%. (c) AcOH, concentrated HCl, aqueous NaNO2, 0–5 °C, aqueous KOAc, malonaldehyde sodium salt (MDA-Na), 0–5 °C, 54%. (d) Methylhydrazine sulfate, KOAc, EtOH, reflux, 74%.
high overlap between the n–π* band of cis-S4 and the long-wavelength absorption band of trans-S4 led to a relatively low cis–trans conversion (85%). In contrast to cis-S4, cis-S3 and cis-S5 exhibited n–π* transitions red-shifted to 25 nm and 36 nm (Table S1†), respectively, which led to partial separation of the n–π* bands of the cis and trans isomers, thereby inducing high (91% S3) to near-quantitative (>95% S5) isomerization.

Theoretical analysis of bidirectional visible-light-driven photoswitches

The geometries of O4, S3, S4, and S5 and their electronic transition characteristics were calculated using density functional theory (DFT) modelling to evaluate the relationship between molecular structures and photophysical properties (see Section 3 in the ESIF†). All trans-isomers exhibited a planar structure with a C–N–N–C dihedral angle of 180.0° (Fig. 2e and S5†), which resulted in symmetry-forbidden n–π* transitions (S0 → S1) with negligible oscillator strength (f = 0.00, Table S4†). Compared to trans-O4, the 4-methylthioarylazopyrazole series showed more effective extension of the π-conjugated system due to the increase of p–π conjugation by introducing the 4-SMe group, as indicated by their frontier molecular orbitals (Fig. S7–S10†). Consequently, the energy gap between the HOMO (π orbital) and LUMO (π* orbital) was smaller for the 4-methylthioarylazopyrazole series (3.714, 3.776, and 3.542 eV for S3, S4, and S5, respectively, as depicted in Fig. 2f), leading to a bathochromic shift of their π–π* transition (S0 → S1). The increased red-shift of S5 was caused by a “complete” conjugation pathway between 5-pyrazole and the azo group that further expands the π-conjugation of the system.40,41 cis-S4 showed a nearly T-shaped conformation with a C–C–N–N dihedral angle of 82.8° (Fig. 2e and S5†), resulting in a weak n–π* transition with f of only 0.0048. In contrast, cis-S3 and cis-S5 were found to disfavor the T-shaped conformation owing to the presence of the ortho nitrogen atom (Fig. 2e and S5†). Therefore, their n–π* absorbance bands were remarkably enhanced (f = 0.1068 and 0.1096 for cis-S3 and cis-S5, respectively, Table S4†), and nearly quantitative conversions from cis to trans isomers were achieved.

Design of low-temperature charging/discharging MOST-PCMs

Thus, all the above data indicate that S5 can be used as a bidirectional visible-light-driven photoswitch, which provided near-quantitative trans–cis and cis–trans photoconversions in

Fig. 3 (a) The structures of An-S5 and Bn-S5. The Tm of trans and cis isomers of (b) An-S5 and (c) Bn-S5. (d) Photographs, optical microscopy images, and POM image of B7-S5 on the ice surface. From top to bottom, the panels are trans-crystals before irradiation, cis-liquids after 400 nm light (40 mW cm−2) irradiation, and regenerated trans-crystals after subsequent irradiation by 532 nm light (110 mW cm−2). The UV-Vis absorption spectra change during (e) 400 nm (40 mW cm−2) and (f) 532 nm (110 mW cm−2) light irradiation of neat B7-S5 at −1 °C. (g) XRD pattern change of B7-S5 on glass slides during the phase transition at −1 °C. (h) Crystal packing structure of trans-B7-S5.
acetonitrile solution. However, pristine S5 was not able to store both visible-light energy and phase transition latent heat at low temperature since the photoisomerization from trans-crystals to cis-liquid was inhibited even at room temperature. To charge and discharge energy below 0 °C, two additional principles are considered: (i) the $T_m$ of the cis-isomer should be below 0 °C, thus forming amorphous cis-liqids to store photon energy and ambient heat; (ii) the trans-isomer should have a $T_m$ and crystallization point ($T_{cr}$) much higher than 0 °C, thus forming trans-crystals to release energy. In order to adjust the $T_m$ of the trans and cis isomers, we varied the length of linear alkyl chains with or without a vinylic end group on the thioalkyl group of S5, denoted as An-S5 and Bn-S5 (Fig. 3a and Scheme S1†). Studying the phase behaviour of the systems using differential scanning calorimetry (DSC), we find that the trans-isomers with longer and intermediate alkyl chain lengths ($n = 6$–12 for An-S5, and $n = 7$, 9, and 11 for Bn-S5) showed $T_m$ (40–60 °C) and $T_{cr}$ above 0 °C (10–45 °C, Tables S5 and S6†). The cis-isomers with intermediate alkyl chain lengths had the lowest $T_m$ (5 °C for A6-S5 and $–1$ °C for B7-S5), as shown in Fig. 3b, c and S13, S14.† Hence, B7-S5 could undergo a reversible visible-light-triggered trans-crystal ↔ cis-liquid transition in cold environments thanks to its low cis-isomer $T_m$ (below 0 °C) and high trans-isomer $T_{cr}$ (34 °C). This property illustrates, to the best of our knowledge, the first example of a functional low-temperature visible-light controlled energy storage MOST-PCM system.

**Charging/discharging below 0 °C**

The charging and discharging processes of B7-S5 at $–1$ °C are discussed as follows. As shown in Fig. 3d and Video S1,† after 400 nm light (40 mW cm$^{-2}$) irradiation, trans-B7-S5 in the orange crystal state lost birefringence and melted into red liquids, which then switched back to a crystal state by 532 nm light (110 mW cm$^{-2}$) irradiation. UV-Vis spectroscopy was used to record the photoisomerization yields during irradiation of a neat sample of B7-S5 with a mass of 2 mg and a thickness of about 50 μm. During the charging process (Fig. 3e), the trans-cis isomerization of B7-S5 in a neat state proceeded easily and produced a high yield of photoisomerization (93%) at 40 min, slightly lower than that in dilute solutions (>95%). During the discharging process (Fig. 3f), the trans-isomer content of the sample increased exponentially, and a near-quantitative (95%) cis-trans isomerization was achieved within 2 min of irradiation. X-ray diffraction (XRD) analyses were carried out to further understand and verify the reversible trans-crystal ↔ cis-liquid behavior of B7-S5 (Fig. 3g). The sharp peaks at 2θ of 5–35° before irradiation corresponded to the regular stack of azo molecules in trans-crystals. After exposure to 400 nm light (40 mW cm$^{-2}$), the peaks disappeared, indicating that cis-B7-S5 had an amorphous structure. Subsequently, XRD patterns were recovered when the sample was irradiated with 532 nm light (110 mW cm$^{-2}$). The X-ray crystal structure of trans-B7-S5 reveals an antiparallel packing, and several weak contacts such as alkyl-alkyl, alkyl-phenyl, and alkyl-pyrazolyl dominating the intermolecular interactions (Fig. 3e). Presumably, the absence of strong intermolecular interactions in trans-B7-S5 offered enough flexibility to the system, which was beneficial for the charging process.42,43

**Energy storage time and energy density**

The cis-isomers of azo molecules can thermally relax into trans-isomers in the dark spontaneously, which determines the energy storage stability of cis-B7-S5. The change of absorbance value at $\pi-\pi^*$ $\lambda_{max}$ (385 nm) as a function of time was measured in acetonitrile solution between 25 and 40 °C (Fig. 4a and S15†). The thermal isomerization rate constants, $k_{cis-\rightarrow trans}$ (Table S8†), were calculated based on the first-order reaction kinetics at 25 °C, 30 °C, 35 °C, and 40 °C, respectively. Based on the Arrhenius equation, cis-B7-S5 was found to have a half-life $t_{1/2}$ of 22.4 days at 0 °C, indicating its stable thermal energy storage capacity at low temperatures. Thermal cis → trans kinetics was also studied in neat states (Fig. S16 and Table S9†), and cis-B7-S5 liquid still had a $t_{1/2}$ as long as 6.3 days at 0 °C. It is ubiquitous to observe a lower $t_{1/2}$ of photoswitches in condensed states than in solution. Cis-isomers tend to adopt fully relaxed geometries in solution, whereas in condensed states different geometries may be preferred due to intermolecular interactions, characterized by lower isomerization barriers.44 Additionally, B7-S5 demonstrated excellent photon-harvesting ability with a quantum yield $\Phi_{trans-\rightarrow cis}$ of 0.39 ± 0.01 for photoisomerization in acetonitrile solution (Fig. 4b and S17†), similar to other reported azopyrazoles compounds.19,41,45

To evaluate the energy density of the MOST-PCM, the isomerization enthalpy $\Delta H_{cr}$ of the thermally induced cis-liquid to trans-liquid reversion reaction and crystallization enthalpy $\Delta H_{cry}$ of the trans-liquid to trans-crystal transition were measured by DSC. As shown in Fig. 4c, the cis-B7-S5 liquid revealed a broad exothermic peak over 60–120 °C during the thermally activated cis-trans isomerization, and the integrated area under the peak represented a $\Delta H_{iso}$ of 0.14 MJ kg$^{-1}$ (44 kJ mol$^{-1}$). This result was consistent with our calculations (49 kJ mol$^{-1}$, Table S7†) based on DFT and similar to other reported azobenzene (<50 kJ mol$^{-1}$). Furthermore, the DSC cooling curve displayed a sharp exothermic peak at around 33 °C with a $\Delta H_{cry}$ of 0.11 MJ kg$^{-1}$ (35 kJ mol$^{-1}$), which was due to the trans-liquid to trans-crystal transition. Therefore, the total thermal energy density of the MOST-PCM was 0.25 MJ kg$^{-1}$ (79 kJ mol$^{-1}$). According to $\Phi_{trans-\rightarrow cis}$ and $\Delta H_{iso}$, the solar efficiency $\eta$ was estimated to be up to 1.3% (see Section 6 in the ESI†), which was one of the highest values reported for azo-based MOST systems (0.2–1.3%).19,28

**Rechargeable energy storage coatings**

To illustrate the practical applications of B7-S5, trans-B7-S5 and diethoxymethylsilane modified chain-like silica were co-dissolved in ethanol and isopropanol (1 : 1, v/v) solution, and then the mixture was drop-cast onto a glass substrate to form rechargeable coatings, as shown in Fig. 5a. The modified chain-like silica formed a transparent porous network structure46 on glass substrates to prevent the leakage of cis-liquid (Fig. S18†). Various patterns were created on the same rechargeable glass substrate through selectively writing/erasing processes (Fig. 5b).
First, a mask was placed on the glass and irradiated with 400 nm blue light. The exposed area of the glass was transformed from the initial opacity to semi-transparency, and the color changed from orange to red. As a result, the target pattern was written on the glass substrate. Subsequently, the erasing process could happen when the substrate without a mask was exposed to 400 nm blue light irradiation, resulting in a globally semi-transparent glass. And then, this glass substrate could be further patterned via selectively irradiating with 532 nm green light. Finally, the glass recovered to its opaque state after exposure to 532 nm green light without a mask. The transmittance spectra of the rechargeable glass under different light irradiations were recorded (Fig. 5c). The glass sheet in a discharged state had a low transmittance of less than 10% in the wavelength range of 500–800 nm, and after exposure to 400 nm blue light, the glass sheet is charged with the transmittance up to ~70% in the wavelength range of 650–800 nm. In addition, the rechargeable glass showed good durability under alternating blue light and green light irradiations (Fig. 5d).

Thanks to the high transmittance of the charged state glass sheet and the low-temperature phase transition of B7-S5, the rechargeable glass had the potential for use as a photochromic solar thermal energy storage window in daily life, especially in cold winters. As a proof-of-principle study, the rechargeable glass sheets were installed on a miniature house model as windows (size 10 by 12 mm and coating thickness ~400 μm). The house model was placed in ~5°C surroundings to ensure that the surface temperature of the windows was around ~1°C. As shown in Fig. 5e, upon exposure to 400 nm blue light (40 mW cm⁻²), the window stored visible light energy and low-temperature ambient heat while transforming from opacity to semi-transparency. Then, by triggering it with 532 nm green light (110 mW cm⁻²), the stored energy was rapidly released on demand as high-temperature heat.

A high-resolution infrared thermal imaging camera was used to track the temperature changes of the window when exposed to 400 nm (40 mW cm⁻²) and 532 nm light (110 mW cm⁻²) (Fig. 5f, S19 and Videos S2–S5). During the 400 nm light irradiation (charging process), the window exhibited a temperature difference of about 3°C above the ambient temperature, indicating a weak photothermal effect. The charged window reached 21.7°C at 60 s during the 532 nm light irradiation (discharging process), about 22.7°C higher than the cold surroundings. A control experiment of irradiating the discharged window with 532 nm light showed a much lower temperature change (9.3°C), which means that the temperature change between the charged window and environment was mainly due to cis-trans isomerization. Assuming that the cis-trans isomerization was fully completed at approximately 60 s, the corresponding thermal power output was estimated to be 256.2 W m⁻². Such high-temperature heat release also means that the B7-S5 molecules on the surface of the window can act as a photon-driven molecular heat pump, upgrading thermal energy from low to high temperature. Furthermore, the optically controlled heat release makes it possible to reach about an order of magnitude higher temperature gradients than is possible with traditional MOST window coating concepts. These solar thermal energy storage coatings...
show unprecedented performances, including visible-light trigger/storage, high energy density, and recyclable ice-cold charging/discharging, thus holding great promise for future energy management systems.

Conclusions

In conclusion, we have successfully designed a series of bidirectional visible-light switching azo molecules and applied them as MOST-PCMs for storing and releasing solar energy below 0 °C. The molecular design strategies are summarized as follows: (i) the 4-thioalkyl substituent on azo molecules shifts the π–π* absorption bands to long-wavelength, enabling bidirectional visible light photoisomerization; (ii) replacing one phenyl ring on the azo molecules with a pyrazole ring increases the half-life of the metastable cis-isomer; (iii) varying the length of thioalkyl chains changes the intermolecular forces, which could adjust the T_m of both trans and cis isomers. Eventually, reversible visible-light-triggered trans-crystal ↔ cis-liquid transitions are achieved below 0 °C. Accordingly, the azo molecules can simultaneously store visible-light energy and low-temperature ambient heat to achieve a high energy density (0.25 MJ kg⁻¹).

Moreover, a rechargeable coating is prepared by drop-coating a solution containing azo photoswitches and modified chain-like silica on the glass surface. The coating shows potential as energy storage windows due to optical transmittance in the charged state, but it is also clear that more work is needed to increase the optical transmittance of the material. Future studies could focus on red-shifting λ_max of azo molecules to the near-infrared region to fabricate efficient semitransparent energy storage windows. Other possible application areas are functional coatings and fabrics with controllable heat release functions. We note that the structure–property relations derived from the chemical design provide a blueprint for how to design future MOST-PCM systems with tailored temperature functions and optimised optical properties. We envision that this work can open an avenue for the design of advanced MOST-PCM systems that store natural sunlight and ambient heat over a wide temperature range.

Data availability

All experimental procedures and data related to this study can be found in the ESI.†

Author contributions

T. L. and K. M. P. conceived the study and supervised the project. Z. S. was primarily responsible for the experiments. W.
S., Z.-Y. Z. and D. F. helped synthesize the compounds. S. W., R.-Z. W. and T.-X. L. provided the infrared thermal imaging camera characterization and analysis. Z.-Y. Z., Z. W. and C. D. contributed to the visualization. X. H. performed the XRD data analysis. Y. H. contributed to DFT calculations. All authors contributed to the writing of the manuscript.

Conflicts of interest
The authors declare no competing interests.

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Notes and references

1. Y. Peng and Y. Cui, Joule, 2020, 4, 724–742.
2. T. Du, Z. Xiong, L. Delgado, W. Liao, J. Peoples, R. Kantharaj, P. R. Chowdhury, A. Marconnet and X. Ruan, Nat. Commun., 2021, 12, 4915.
3. R. Hu, Y. Liu, S. Shin, S. Huang, X. Ren, W. Shu, J. Cheng, G. Tao, W. Xu, R. Chen and X. Luo, Adv. Energy Mater., 2020, 10, 1903921.
4. A. L. Moore and L. Shi, Mater. Today, 2014, 17, 163–174.
5. D. Li, J. Wang, Y. Ding, H. Yao and Y. Huang, Appl. Energy, 2019, 236, 1168–1182.
6. T. R. Whiffen and S. B. Riffat, Int. J. Low-Carbon Technol., 2013, 8, 159–164.
7. A. Sasic Kalagasidis, Journal of Building Performance Simulation, 2014, 7, 289–308.
8. M. Song, F. Niu, N. Mao, Y. Hu and S. Deng, Energy Build., 2018, 158, 776–793.
9. G. G. D. Han, H. Li and J. C. Grossman, Nat. Commun., 2017, 8, 1446.
10. Z. Wang, P. Erhart, T. Li, Z.-Y. Zhang, D. Sampredo, Z. Hu, H. A. Wegner, O. Brummel, J. Libuda, M. B. Nielsen and K. Moth-Poulsen, Joule, 2021, 5, 3116–3136.
11. T. J. Kucharski, N. Ferralis, A. M. Kolpak, J. O. Zheng, D. G. Nocera and J. C. Grossman, Nat. Chem., 2014, 6, 441–447.
12. Q. Qiu, Y. Shi and G. G. D. Han, J. Mater. Chem. C, 2021, 9, 11444–11463.
13. M. Quant, A. E. Hillers-Bendtsen, S. Ghasemi, M. Erdelyi, Z. Wang, L. M. Muhammad, N. Kann, K. V. Mikkelsen and K. Moth-Poulsen, Chem. Sci., 2022, 13, 834–841.
14. M. Manso, A. U. Petersen, Z. Wang, P. Erhart, M. B. Nielsen and K. Moth-Poulsen, Nat. Commun., 2018, 9, 1945.
15. F.-Y. Meng, I. H. Chen, J.-Y. Shen, K.-H. Chang, T.-C. Chou, Y.-A. Chen, Y.-T. Chen, C.-L. Chen and P.-T. Chou, Nat. Commun., 2022, 13, 797.
16. L. Dong, Y. Feng, L. Wang and W. Feng, Chem. Soc. Rev., 2018, 47, 7339–7368.
17. H. Liu, J. Tang, L. Dong, H. Wang, T. Xu, W. Gao, F. Zhai, Y. Feng and W. Feng, Adv. Funct. Mater., 2021, 31, 2008496.
18. K. Ishiba, M.-a. Morikawa, C. Chikara, T. Yamada, K. Iwase, M. Kawakita and N. Kimizuka, Angew. Chem., Int. Ed., 2015, 54, 1532–1536.
19. Z.-Y. Zhang, Y. He, Z. Wang, J. Xu, M. Xie, P. Tao, D. Ji, K. Moth-Poulsen and T. Li, J. Am. Chem. Soc., 2020, 142, 12256–12264.
20. M. A. Gerkman, R. S. L. Gibson, J. Calbo, Y. Shi, M. J. Fuchter and G. G. D. Han, J. Am. Chem. Soc., 2020, 142, 8686–8695.
21. X. Xu and G. Wang, Small, 2022, 2107473.
22. L. Fei, Y. Yin, M. Yang, S. Zhang and C. Wang, Energy Storage Mater., 2021, 42, 636–644.
23. C. M. Z. Samuel, W. L. Shames, N. Ferralis and J. C. Grossman, International Journal of Energy Efficient Design, 2013, 15, 1114–1119.
24. O. S. Bushuyev, T. C. Corkery, C. J. Barrett and T. Frišičić, Chem. Sci., 2014, 5, 3158–3164.
25. O. S. Bushuyev, A. Tomberg, T. Frišičić and C. J. Barrett, J. Am. Chem. Soc., 2013, 135, 12556–12559.
26. W.-C. Xu, S. Sun and S. Wu, Angew. Chem., Int. Ed., 2019, 58, 9712–9740.
27. A. K. Saydjari, P. Weis and S. Wu, Adv. Energy Mater., 2017, 7, 1601622.
28. Y. Shi, M. A. Gerkman, Q. Qiu, S. Zhang and G. G. D. Han, J. Mater. Chem. A, 2021, 9, 9798–9808.
29. A. A. Beharry, O. Sadowski and G. A. Woolley, J. Am. Chem. Soc., 2011, 133, 19684–19687.
30. D. Bléger, J. Schwarz, A. M. Brouwer and S. Hecht, J. Am. Chem. Soc., 2012, 134, 20597–20600.
31. L. Fei, Y. Yin, J. Zhang and C. Wang, Sol. RRL, 2020, 4, 2000499.
32. Z. Refaa, A. Hofmann, M. F. Castro, J. O. Hernandez, Z. Wang, H. Hölzel, J. W. Andreasen, K. Moth-Poulsen and A. S. Kalagasidis, Appl. Energy, 2022, 310, 118541.
33. X. Yang, G. Ma, S. Zheng, X. Qin, X. Li, L. Du, Y. Wang, Y. Zhou and M. Li, J. Am. Chem. Soc., 2020, 142, 9460–9470.
34. R. J. Tombari, J. R. Tuck, N. Yardeny, P. W. Gingrich, D. J. Tantillo and D. E. Olson, Org. Biomol. Chem., 2021, 19, 7575–7580.
35. Z.-Y. Zhang, Y. He, Y. Zhou, C. Yu, L. Han and T. Li, Chem.– Eur. J., 2019, 25, 13402–13410.
36. J. A. Pople, Angew. Chem., Int. Ed., 1999, 38, 1894–1902.
37. S. Bhunia, A. Dolai and S. Samanta, Chem. Commun., 2020, 56, 10247–10250.
38. X. Xu, B. Wu, P. Zhang, Y. Xing, K. Shi, W. Fang, H. Yu and G. Wang, ACS Appl. Mater. Interfaces, 2021, 13, 22665–22663.
39. D. Fang, Z.-Y. Zhang, Z. Shangguan, Y. He, C. Yu and T. Li, J. Am. Chem. Soc., 2021, 143, 14502–14510.
40. Y. He, Z. Shangguan, Z.-Y. Zhang, M. Xie, C. Yu and T. Li, Angew. Chem., Int. Ed., 2021, 60, 16539–16546.
41. J. Calbo, C. E. Weston, A. J. P. White, H. S. Rzepa, J. Contreras-Garcia and M. J. Fuchter, J. Am. Chem. Soc., 2017, 139, 1261–1274.
42 P. Kumar, A. Srivastava, C. Sah, S. Devi and S. Venkataramani, Chem.–Eur. J., 2019, 25, 11924–11932.
43 L. Kortekaas, J. Simke, D. W. Kurka and B. J. Ravoo, ACS Appl. Mater. Interfaces, 2020, 12, 32054–32060.
44 K. Janus and J. Sworakowski, J. Phys. Chem. B, 2005, 109, 93–101.
45 C. E. Weston, R. D. Richardson, P. R. Haycock, A. J. P. White and M. J. Fuchter, J. Am. Chem. Soc., 2014, 136, 11878–11881.
46 Y. Li, Z. Zhang, M. Wang, X. Men and Q. Xue, J. Mater. Chem. A, 2017, 5, 20277–20288.