Warm/cool-tone switchable thermochromic material for smart windows by orthogonally integrating properties of pillar[6]arene and ferrocene

Sai Wang¹, Zuqiang Xu², Tingting Wang², Tangxin Xiao³, Xiao-Yu Hu², Ying-Zhong Shen¹ & Leyong Wang²,³

Functional materials play a vital role in the fabrication of smart windows, which can provide a more comfortable indoor environment for humans to enjoy a better lifestyle. Traditional materials for smart windows tend to possess only a single functionality with the purpose of regulating the input of solar energy. However, different color tones also have great influences on human emotions. Herein, a strategy for orthogonal integration of different properties is proposed, namely the thermo-responsiveness of ethylene glycol-modified pillar[6]arene (EGP6) and the redox-induced reversible color switching of ferrocene/ferrocenium groups are orthogonally integrated into one system. This gives rise to a material with cooperative and non-interfering dual functions, featuring both thermochromism and warm/cool tone-switchability. Consequently, the obtained bifunctional material for fabricating smart windows can not only regulate the input of solar energy but also can provide a more comfortable color tone to improve the feelings and emotions of people in indoor environments.
Smart windows have great potential as a way of improving building energy efficiency and indoor comfort, by reversibly modulating the amount of visible light and solar radiation entering into the buildings\textsuperscript{1–3}. To achieve switchable optical performance of such smart windows, avoiding unnecessary energy consumption and unsatisfactory indoor environment, a large amount of prior research has been performed, which mostly focused on controlling the transmittance of windows by external stimuli, including electricity (electrochromism)\textsuperscript{4–10}, light (photochromism)\textsuperscript{11–13}, heat (thermochromism)\textsuperscript{14–16}, and gas (gasochromism)\textsuperscript{17}. Amongst these, thermochromic smart windows have attracted specific attention because they can automatically adjust the window transparency through weather-responsive control without extra energy input. This results in a reversible regulation of the solar transmittance (mainly in the near-infrared region), automatically adjusting the amount of solar heat entering into the buildings\textsuperscript{18–20}. So far, vanadium dioxide (VO\textsubscript{2}) is one of the most widely used inorganic materials for the fabrication of thermochromic smart windows, but the application of VO\textsubscript{2}-based smart windows is limited by many shortcomings, such as a high transition temperature (68 °C), low light transmittance, and unattractive blackness\textsuperscript{21–23}. Meanwhile, to the best of our knowledge, few studies have achieved the construction of smart windows based on organic materials, and most of the known systems are limited to a regulation of solar energy input, thus only possessing a single functionality\textsuperscript{24–30}.

However, taking into account emotional and psychological effects, not only the amount of light input, but also the color of the light has a tremendous importance. In color science, color can be categorized into warm color, cool color, and neutral color\textsuperscript{31,32}. Warm-tone colors, such as red, orange, and yellow makes people feel warm, lively, and excited, while cool-tone colors including green, blue, and purple gives us a feeling of coolness, serenity, and calmness. Smart windows that are capable of providing warm colors in winter and cool colors in summer will not only make our homes much more comfortable, but can also be beneficial to improve human emotions. Although, thermochromic supramolecular materials based on charge-transfer interactions with excellent color switchable property developed by Fang and Olson have attracted great interest and attention\textsuperscript{33,34}, their application in the fabrication of smart windows has yet to be realized. Accordingly, developing an ideal warm/cool tone switchable thermochromic material for fabricating smart windows is an amazing and challenging endeavour, which could have an immediate positive effect on improving the quality of life for people throughout the world.

Orthogonal self-assembly refers to an integration strategy of two or more types of non-covalent interactions with high specificity and selectivity\textsuperscript{35–42}. Using this orthogonal strategy, various types of supramolecular complexes (such as ligand-receptor, host–guest) have been applied for the fabrication of smart and functional materials based on supramolecular polymers. However, up to now, most orthogonal self-assembly strategies rely on selective orthogonal supramolecular interactions, while the individual functional properties of the interacting entities are not fully explored. For example, pillar[n]arenes\textsuperscript{43–53}, as a family of macrocyclic hosts, not only possess outstanding properties in host–guest chemistry, but can also undergo facile modification to introduce additional fascinating functionalities, such as hydrophobicity for super-hydrophobic surfaces\textsuperscript{54}, water-solubility\textsuperscript{55}, biofilm inhibition\textsuperscript{47}, multi-responsiveness to CO\textsubscript{2}\textsuperscript{56}, light\textsuperscript{48}, temperature\textsuperscript{57}, and so on. Inspired by the concept of orthogonal self-assembly, we try to orthogonally integrate the unique properties of different supramolecular substrates to achieve the fabrication of smart materials with cooperative and non-interfering functions.

Herein, a dry gel with ferrocene pendants modifying on the polycrylamide-based polymer networks (Fc-gel) has been successfully prepared. A well-swollen and high transparent hydrogel (Fc-gel-EGP6) is obtained when the Fc-gel is immersed in the

**Fig. 1** Chemical structures and schematic illustration of the warm/cool tone switchable thermochromic material for smart windows. Preparation of Fc-gel+EGP6 hydrogel and the schematic illustration of warm/cool tone switchable thermochromic material based on the orthogonal integration strategy of the unique properties of pillar[6]arene and ferrocene.
aqueous solution of ethylene glycol chain-modified pillar[6]arene (EGP6). This phenomenon is due to the formation of hydrophilic inclusion complexes between EGP6 and the ferrocene groups of Fc-gel instead of the original hydrophobic domains formed by physical cross-linking of the hydrophobic ferrocene units. Thus, water absorption capacity of the generated Fc-gel·EGP6 hydrogel is significantly improved, which results in a drastically improved swelling behaviour and an enhanced transparency of the Fc-gel·EGP6 hydrogel in comparison to the dry Fc-gel. Moreover, owing to the thermo-responsive properties of EGP6 and the reversible transformation between orange and green colors of ferrocene/ferrocenium groups under redox-response, we have orthogonally integrated the functional properties of both supramolecular building blocks based on host–guest interaction in order to design a warm/cool tone-switchable thermochromic material. This material is highly suitable for the fabrication of smart windows with dual functionality, both for regulating the input of solar energy and for improving the feelings and emotions of inhabitants for a comfortable life (Fig. 1).

Results
Preparing Fc-gel·EGP6 hydrogel by host–guest complexation.
Initially, the complexation between EGP6 and Fc-containing polymer networks was investigated by $^1$H NMR spectroscopy. Since both the polymer networks and the hydrophobic ferrocene moiety are insoluble in water, a water-soluble linear polymer mPEG-Fc was selected as a model guest for the host–guest complexation study (Fig. 2 and Supplementary Fig. 2). When a mixture of EGP6 and mPEG-Fc with a 1:1 molar ratio was dissolved in D$_2$O, the peaks for protons on EGP6 were shifted to low field, with the most significant downfield chemical shift observed for the aromatic protons $H_A$ (Δ$\delta$ = 0.14 ppm). Meanwhile, the signal for the ferrocene-protons ($H_B$, including one methylene unit) of mPEG-Fc showed an obvious upfield chemical shift (overlapped with the peaks of EGP6) due to the shielding effect of the electron-rich cavity of EGP6. The above results reveal that the ferrocene moiety of model guest mPEG-Fc is threaded into the hydrophobic cavity of EGP6. This was also confirmed by a 2D NOESY experiment (Supplementary Figs. 4 and 5), which clearly showed the NOE correlation signals between protons $H_A$ of EGP6 and part of protons $H_B$ on the ferrocene moiety of mPEG-Fc.

Subsequently, a Job’s plot analysis based on UV–vis spectra data showed that the complexation between EGP6 and the ferrocene moiety of mPEG-Fc had 1:1 stoichiometry (Supplementary Fig. 1). The corresponding binding constant of EGP6 $\supset$ mPEG-Fc in water was determined to be $(3.98 \pm 0.10) \times 10^2$ M$^{-1}$ by a non-linear curve-fitting method (Supplementary Fig. 3). In this case, hydrophobic interaction is the main driving force for such a host–guest complexation, leading to the formation of a stable 1:1 inclusion complex between EGP6 and the ferrocene moiety.

Having demonstrated the Fc-EGP6 interaction in the model substrate, we turned our attention to the Fc-substituted polymer networks for the following investigation. A disc-shaped sample consisting of cross-linked random copolymers (Fc-gel) was prepared by free radical copolymerization of acrylamide, a ferrocene-modified monomer, and NN’-methylenebis(acrylamide) (MBA) under conventional conditions (for details, see preparation of Fc-gel in Supplementary Discussion). Based on our previous work $^{55}$, the obtained Fc-gel should display a significant swelling behavior in an aqueous solution of water-soluble pillar[6]arene. The swelling ratio will be dramatically promoted with an increasing amount of the incorporated ferrocene subunits from 0% to 10 mmol%; whereas, when the ferrocene loading was more than 10 mmol%, a decreased swelling ratio is expected induced by the dominant hydrophobic effect of the ferrocene groups. Therefore, in this study, the molar ratio of ferrocene subunits in Fc-gel was selected as 10 mmol% for sufficient host–guest complexation with EGP6 and subsequent excellent swelling behavior of Fc-gel.

The samples of Fc-gel were immersed in water or EGP6 aqueous solution at 25 °C to investigate their different swelling behaviors. When immersed in EGP6 aqueous solution (15 mM),
the dry Fc-gel showed dramatic swelling, and the diameter of the disc increased from 7 to 17.5 mm with an increment of 150% compared to the dried sample (Fig. 3c and Supplementary Fig. 6). Moreover, the obtained sufficiently swollen hydrogel exhibited a homogeneous morphology with a porous network structure, as observed from scanning electron microscopy (SEM) image (Fig. 3b). In contrast, when the dry Fc-gel was immersed in pure water as a control experiment, the equilibrium diameter only increased by 64.3% (from 7 to 11.5 mm, Fig. 3c and Supplementary Fig. 6), and the obtained hydrogel exhibited a heterogeneous morphology with denser matrix and a non-uniform porous structures observed from the SEM image (Fig. 3a). The main reason for this different swelling behavior can be attributed to the formation of hydrophilic EGP6-ferrocene inclusion complexes, which transformed the hydrophobic ferrocene groups into hydrophilic moieties and resulted in the obviously improved water absorption capacity and drastically increased swelling property of the Fc-gel in the presence of EGP6.

Fabricating orthogonal functional material for smart windows.

The response to external stimuli is a significant property of smart hydrogels. Since EGP6 shows a unique thermo-responsiveness and the ferrocene moiety exhibits a reversible transformation between orange and green ferrocene/ferrocenium groups under redox-stimuli, we envisaged the orthogonal integration of both of the above unique properties to design a warm/cool tone-switchable thermochromic material for fabricating smart windows. Initially, we investigated the stimuli-responsive properties of EGP6 and ferrocene in aqueous solution. When EGP6 aqueous solution was heated up to 40 °C, a white and turbid solution was obtained, which could return to a clear solution after being cooled to 25 °C (Fig. 4a, b). Meanwhile, mPEG-Fc in its aqueous solution exhibited an orange color. Whereas, upon oxidation with ammonium persulfate, the original orange solution gradually became dark green due to the transformation of ferrocene into ferrocenium (Fc⁺), which could be reversed by subsequent reduction process, restoring the original orange color (Fig. 4c, d, for the host–guest interaction between Fc⁺ and EGP6, see Supplementary Figs. 8–10). Based on these observations, we tried to orthogonally integrate the above unique properties of EGP6 and ferrocene to the obtained hydrogel for further fabricating of a warm/cool tone-switchable thermochromic material for smart windows.

**Fig. 3** SEM micrographs and photographs of hydrogels. SEM micrographs of freeze-dried Fc-gel immersed in a pure water and b EGP6 aqueous solution (15 mM); Scale bars are 50 µm. c the corresponding photographs of Fc-gel under different conditions.
For the construction of smart windows, transparency is one of the most important prerequisites to ensure indoor illumination and input of solar energy. Therefore, the prepared functional materials for smart windows should firstly be transparent. The obtained well-swollen Fc-gel-EGP6 hydrogel shows a high grade of transparency and an orange color, thus providing an excellent foundation for its potential application in constructing windows (Fig. 3c). Investigation of the optical transparency of Fc-gel-EGP6 hydrogel by UV–vis transmission spectrum (Supplementary Fig. 7) showed enhanced transmittance with increasing wavelength in the visible region (380–780 nm) at 25 °C. More importantly, the hydrogel showed a high transmittance from 79.4 to 89.6% in the near infrared region (780–1100 nm). Thus, the Fc-gel-EGP6 hydrogel is ideally suited to allow an excellent input of solar energy, effectively increase the indoor temperature with windows prepared by such a functional material. Moreover, the orange warm-toned color of the material makes it a potential candidate for smart windows with the ability to make indoor people feel warm and lively.

We next turned our attention to a possible thermo-responsiveness of the material. From an application point-of-view, we aspired to minimize the input of solar energy when the indoor temperature was too high to live comfortably. Especially in summer, thermo-responsive windows would represent an effective means of reducing the indoor temperature and thus decrease the electrical energy consumption of air conditioning devices. Therefore, we can confirm that the transparency/opaqueness of the Fc-gel-EGP6 hydrogel is reversible and highly reproducible for many alternating cycles between 25 and 40 °C. With regard to the second functionality of the material, the warm/cool tone color change, we subsequently investigated its reversible redox-behaviour. When the Fc-gel-EGP6 hydrogel was immersed in ammonium persulphate aqueous solution ((NH4)2S2O8), the gel gradually changed its color from orange to green within 8 min (Fig. 5c and Supplementary Movie2). This phenomenon is based on the thermal-responsiveness of EGP6. At low temperatures, the Fc-gel-EGP6 hydrogel will be formed driven by host-guest interaction. However, upon increasing the temperatures above its cloud point temperature (Tcloud), EGP6 will aggregate and microseparate from water inside the hydrogel due to the interaction of the hydrophobic groups of pillararene backbone37, and finally, resulting in the generation of an opaque Fc-gel-EGP6 hydrogel with the ability to prevent visible light and solar radiation from passing through. When decreasing the temperature from 40 to 25 °C, the opaque Fc-gel-EGP6 hydrogel immediately returned to its original transparency because of the re-dissolution of EGP6 below its Tcloud (Supplementary Movie 1). Meanwhile, such reversible thermo-responsive behavior was also supported by variable-temperature 1H-NMR spectroscopy (Supplementary Fig. 11). More importantly, after more than 100 cycles of alternating temperature increasing/decreasing, the transmittances kept almost constant between two values measured at 1099 nm (Fig. 7a). In addition, the Fc-gel-EGP6 hydrogel remained very good stability without any leakage of EGP6 from the hydrogel backbone even after 100 thermal cycles (Supplementary Fig. 12). Therefore, we can confirm that the transparency/opaqueness of the Fc-gel-EGP6 hydrogel is reversible and highly reproducible for many alternating cycles between 25 and 40 °C.

With regard to the second functionality of the material, the warm/cool tone color change, we subsequently investigated its reversible redox-behaviour. When the Fc-gel-EGP6 hydrogel was immersed in ammonium persulphate aqueous solution ((NH4)2S2O8), the gel gradually changed its color from orange to green within 8 min (Fig. 5c and Supplementary Movie 2). This is mainly due to the fact that the original color of the hydrogel was contributed by the orange ferrocene group, which is transformed into green ferrocenium moiety upon oxidation (Fig. 4c, d). To our delight, the oxidation process had hardly any impact on the transmittance of the Fc-gel-EGP6 hydrogel and the obtained oxidized hydrogel with green color also exhibited high transparency. From the UV–vis transmission spectra (Fig. 6c), a small absorption peak at about 620 nm could be observed, giving direct spectroscopic evidence of the ferrocenium-moiety. Meanwhile, similar as the Fc-gel-EGP6 hydrogel, the oxidized hydrogel...
with dark green color also had excellent thermochromism (Figs. 5c and 6d, and Supplementary Movie 3), and the reversible transformation between transparency and opaqueness of the oxidized hydrogel were also highly reproducible for more than 100 thermal cycles (Fig. 7b). Interestingly, a reverse process for transforming the green hydrogel into its original orange color could be achieved by immersing the oxidized hydrogel in hydrazine hydrate (N₂H₄·H₂O) aqueous solution (Supplementary Movie 2), leading to the reduction of green ferrocenium into orange ferrocene groups. And the regenerated orange hydrogel also maintained its high transparency and good performance of thermochromism (Fig. 7d). Moreover, the above switchable process between orange and green colors exhibited good reversibility and high repeatability based on the oxidation/reduction of the ferrocene groups within the Fc-gel·EGP6 hydrogel (Fig. 7c). More importantly, the oxidized green Fc-gel·EGP6 hydrogel material possesses the characteristics of cool-toned colors. Thus, this material could be used to fabricate smart windows for providing cool and calm feelings of indoor people.

In addition, for the real-life applications of smart windows, a feasible and practical way for affecting the desired switching process is necessary. In this Fc-gel·EGP6 hydrogel system, thermochromic performance depends on the environmental temperature, while the color switching between warm and cool tone can be controlled by a feasible redox stimuli. According to previous work, cyclic voltammetry (CV) study indicated that the pillararene–ferrocene inclusion complexes have stable oxidation and reduction peaks. Therefore, we have fabricated an electrochemical device based on the Fc-gel·EGP6 hydrogel to achieve the color switching between warm and cool tone by applying a suitable electric potential instead of using chemical oxidizing and reducing agents. As shown in Fig. 8, after being immersed in KCl aqueous solution (0.1 M), the Fc-gel·EGP6 hydrogel was sandwiched between two pieces of ITO conductive glass to form a simple device. Upon holding the potential at +4.0 V for 6 min, the color of the material in the device could be completely transformed from orange to green due to the electrochemical oxidation of ferrocene to ferricenium cation. After further holding the potential at –4.0 V for 6 min, the Fc-gel·EGP6 hydrogel was reduced and the color of the material returned to orange. Therefore, based on the orthogonally integrated properties of EGP6 and ferrocene by host–guest interaction, a warm/cool tone-switchable Fc-gel·EGP6 hydrogel was successfully prepared, which could be used as a novel material for fabricating smart windows to improve the emotions of indoor people, meanwhile, both in warm and cool tones, the hydrogel was transparent and possessed excellent thermochromism to control the input of solar energy for a more comfortable indoor environment.

**Discussion**

In summary, we have orthogonally integrated the unique properties of pillar[6]arene and ferrocene based on host–guest interaction to develop a warm/cool tone-switchable thermochromic material for the fabrication of smart windows. Owing to the efficient EGP6-ferrocene complexation, a well-swollen and highly transparent Fc-gel·EGP6 hydrogel was prepared by immersing a ferrocene-modified polyacrylamide-based polymer networks in EGP6 aqueous solution. Moreover, the generated Fc-gel·EGP6 hydrogel exhibited warm/cool tone-switchable property owing to the reversible transformation between orange and green ferrocene/ferricenium groups under redox-stimuli, which has potential applications in fabricating smart windows with specific function for improving the feelings and emotions of indoor people. More importantly, the transparent Fc-gel·EGP6 hydrogel inherited the excellent thermo-responsiveness of EGP6 in both
warm and cool color to become a thermochromic material, which can regulate the input of solar energy for a more comfortable indoor environment. In this study, the unique properties of EGP6 and ferrocene were orthogonally integrated together, where both EGP6 and ferrocene perform its own duty with non-interference but mutual promotion. Therefore, we successfully provide the first applied example for this orthogonal integration strategy of different properties with the purpose of developing a warm/cool tone-switchable thermochromic material for fabricating smart windows, which have promising applications in the field of material science and technology. Furthermore, we believe that this orthogonal integration strategy of different properties might provide a new direction for design and development of novel functional materials.

Methods

General. All reagents were commercially available and used as supplied without further purification unless otherwise stated. EGP6, mPEG-Fc (Mn = 2347 g·mol⁻¹), ferrocene-modified acrylamide monomer (FcAm), and dry Fc-gel were synthesized according to the literatures. Nuclear magnetic resonance (NMR) spectra were recorded on a Bruker Advance DMX 400 spectrophotometer or a Bruker Advance DMX 500 spectrophotometer with internal standard tetramethylsilane (TMS) and solvent signals as internal references at 25 °C. SEM was carried out on a Shimadzu SSX-550 device. UV–vis spectra were recorded on a Shimadzu UV-1780 UV–vis Spectrophotometer.

Preparation method of dry Fc-gel. FcAm (220 mg, 0.70 mmol), acrylamide (448 mg, 6.30 mmol), and N, N'-methylenebis(acrylamide) (MBA) (5.40 mg, 0.035 mmol) were dissolved in dimethyl sulfoxide (DMSO) (1.6 mL). Then azodiisobutyronitrile (AIBN) (28.8 mg, 0.175 mmol) was added to the solution and the mixture was purged with dry argon for 30 min. The resulting solution was equally divided into 10 vials (10 mm in diameter) and sealed. The polymerization was performed in an oven at 70 °C for 24 h. Then, the vials were cooled down to room temperature, disc-shaped samples were washed successively by DMSO and deionized water, and subsequently dried in oven (50 °C) under vacuum for 12 h after

![Graphs and images](image-url)
natural drying. Finally, disc-shaped dry Fe-gel was obtained with 7 mm in diameter and 1 mm in thickness.

Synthesis and characterization. Synthesis and relevant characterization details are provided in the Supplementary Information.

Data availability. The authors declare that the data supporting the findings of this study are available within the paper and its Supplementary Information including Supplementary Methods and Supplementary Discussion. All data are available from the authors on reasonable request.

Received: 14 December 2017 Accepted: 15 March 2018

Published online: 30 April 2018

References

1. Baezens, R., Jelle, B. P. & Gustavsen, A. Properties, requirements and possibilities of smart windows for dynamic daylight and solar energy control in buildings: a state-of-the-art review. Sol. Energy Mater. Sol. Cells 94, 87–105 (2010).

2. Bella, F. et al. A new design paradigm for smart windows: photocurable polymers for quasi-solid photocathodic devices with excellent long-term stability under real outdoor operating conditions. Adv. Funct. Mater. 26, 1127–1137 (2016).

3. Granqvist, C. G. Electrochromics for smart windows: oxide-based thin films and devices. Thin Solid Films 564, 1–38 (2014).

4. Wang, Y., Runnerstrom, E. L. & Milliron, D. J. Switchable materials for smart applications. Chem. Rev. 112, 754–829 (2012).

5. Deb, S. K. Opportunities and challenges in science and technology of WO3 for applications. J. Mater. Chem. A 3, 11515–11588 (2015).

6. Kiruthika, S. & Kulkarni, G. U. Energy efficient hydrogel based smart windows: thin films of tungsten oxide and nickel oxide, and devices based on these. J. Mater. Chem. 17, 127–156 (2007).

7. Thakur, V. K., Ding, G., Ma, J., Lee, P. S. & Lu, X. Hybrid materials and polymers for the construction of diverse electrochromic devices. Adv. Mater. 24, 4071–4096 (2012).

8. Connelly, K., Wu, Y., Chen, J. & Lei, Y. Design and development of a reflective membrane for a novel building integrated concentrating photovoltaic (BICPV) ‘smart window’ system. Appl. Energy 182, 331–339 (2016).

9. Li, N. et al. Interfacial–charge-transfer–induced photochromism of MoO3@TiO2 crystalline–core amorphous–shell nanorods. Sol. Energy Mater. Sol. Cells 163, 231–236 (2017).

10. Wang, Y. et al. Disentangling photochromism and electrochromism by blocking hole transfer at the electrolyte interface. Chem. Mater. 28, 7198–7202 (2016).

11. Lee, H. Y. et al. Thermochromic lignogel: a new class of stimuli responsive materials with super cyclic stability for solar modulation. Chem. Mater. 29, 6947–6955 (2017).

12. Zhou, Y., Cai, Y., Hu, X. & Long, Y. Temperature-responsive hydrogel with ultra-large solar modulation and high luminous transmission for “smart window” applications. J. Mater. Chem. A 2, 13550–13555 (2014).

13. Lin, J. et al. Thermochromic halide perovskite solar cells. Nat. Mater. 17, 261–267 (2018).

14. Feng, W. et al. Gasochromic smart window: optical and thermal properties, energy simulation and feasibility analysis. Sol. Energy Mater. Sol. Cells 144, 316–323 (2016).

15. Chen, Y. et al. High performance and enhanced durability of thermochromic films using VO2@TiO2 core-shell nanoparticles. ACS Appl. Mater. Interfaces 9, 27784–27791 (2017).

16. Granqvist, C. G. Transparent conductors as solar energy materials: a panoramic review. Sol. Energy Mater. Sol. Cells 91, 1529–1598 (2007).

17. Granqvist, C. G., Lansaker, P. C., Miyuka, N. R., Niklasson, G. A. & Avendano, E. Progress in chromogenics: new results for electrochromic and thermochromic materials and devices. Sol. Energy Mater. Sol. Cells 93, 2032–2039 (2009).

18. Gao, Y. et al. Nanocrystalline VO2 thermochromic smart glass: a review on progress in solution processing. Nano Energy 1, 221–246 (2012).

19. Ye, H., Meng, X. & Xu, X. B. Theoretical discussions of perfect window, ideal near infrared solar spectrum regulating window and current thermochromic window. Energ. Build. 49, 164–172 (2012).

20. Zhang, Z. et al. Thermochromic VO2 thin films: solution-based processing, improved optical properties, and lowered phase transition temperature. J. Sol-Gel Sci. Technol. 62, 10738–10747 (2010).

21. Khan, W., Soo, J.-M. & Park, S.-Y. Synthesis and micellization of a novel diblock copolymer of poly(N-isopropylacrylamide)-b-SGCLP and its application in stability of 5CB droplets in aqueous medium. Soft Matter 7, 780–787 (2011).

22. Lee, E., Kim, D. & Yoon, J. Stepwise activation of switchable glazing by compositional gradient of copolymers. ACS Appl. Mater. Interfaces 8, 26359–26364 (2016).

23. Lee, H. Y. et al. A dual-responsive nanocomposite toward climate-adaptable solar modulation for energy-saving smart windows. ACS Appl. Mater. Interfaces 9, 6054–6063 (2017).

24. Rao, M. S. & Dave, B. C. “Smart” glasses: molecular programming of rapid dynamic responses in organosilica sol-gels. Adv. Mater. 14, 443–447 (2002).

25. Zhou, Y., Cai, Y., Hu, X. & Long, Y. VO2/ hydrogel hybrid nanomaterial thermochromic material with ultra-high solar modulation and luminous transmission. J. Mater. Chem. A 3, 1121–1126 (2015).

26. Zhang, K., Zhang, M., Feng, X., Hempensius, M. A. & Vancso, G. J. Switching light transmittance by responsive organometallic poly(ionic liquid): control by cross talk of thermal and redox stimuli. Adv. Funct. Mater. 27, 1702784 (2017).

27. Plass, J. L., Heidig, S., Hayward, E. O., Homer, B. D. & Um, E. Emotional design in multimedia learning: effects of shape and color on affect and learning. J. Edutainment. Instr. 29, 128–140 (2014).

28. Yilmaz, M. D. & Huskens, J. Orthogonal supramolecular interaction motifs for the construction of complex supramolecular architecture. Chem. Soc. Rev. 44, 779–789 (2015).

29. Yuan, T. et al. Tunable thermochromism of multifunctional charge-transfer-based supramolecular materials assembled in water. Chem. Mater. 29, 19937–19945 (2017).

30. Yuan, T. et al. Versatile thermochromic supramolecular materials based on competing charge transfer interactions. Adv. Funct. Mater. 26, 8604–8612 (2016).

31. He, Z., Jiang, W. & Schalley, C. A. Integrative self-sorting: a versatile strategy for the construction of complex supramolecular architecture. Chem. Soc. Rev. 44, 779–789 (2015).

32. Hu, X.-Y., Xiao, T., Lin, C., Huang, F. & Wang, L. Dynamic supramolecular complexes constructed by orthogonal self-assembly. Acc. Chem. Res. 47, 2041–2051 (2014).

33. Kumar, D. K. & Steed, J. W. Supramolecular gel phase crystallization: orthogonal self-assembly under non-equilibrium conditions. Chem. Soc. Rev. 43, 2080–2088 (2014).

34. Li, S.-L., Xiao, T., Lin, C. & Wang, L. Advanced supramolecular polymers constructed by orthogonal self-assembly. Chem. Soc. Rev. 41, 5950–5968 (2012).

35. Wei, P., Yan, X. & Huang, F. Supramolecular polymers constructed by orthogonal self-assembly based on host-guest and metal-ligand interactions. Chem. Soc. Rev. 44, 815–832 (2015).

36. Yan, X. et al. A multiresponsive, shape-persistent, and elastic supramolecular polymer network gel constructed by orthogonal self-assembly. Adv. Mater. 24, 362–369 (2012).

37. Yang, L., Tan, X., Wang, Z. & Zhang, X. Supramolecular polymers: historical development, preparation, characterization, and functions. Chem. Rev. 115, 7196–7239 (2015).

38. Yilmaz, M. D. & Huskens, J. Orthogonal supramolecular interaction motifs for functional monolayer architectures. Soft Matter 8, 11768–11780 (2012).

39. Xue, M., Yang, Y., Chi, X., Zhang, Z. & Huang, F. Pillararenes, a new class of macrocycles for supramolecular chemistry. Acc. Chem. Res. 45, 1294–1308 (2012).

40. Chi, X., Yu, G., Shao, L., Chen, J. & Huang, F. A dual thermo-responsive gemini-type supra-ampiphilic macromolecular [3]pseudorotaxane based on pillar[5]arene/parquat cooperative interaction. J. Am. Chem. Soc. 138, 3168–3174 (2016).

41. Dong, S. et al. LCS-type phase behavior induced by pillar[5]arene/ionic liquid host–guest complexation. Adv. Mater. 25, 8664–8673 (2013).

42. Hu, X.-Y. et al. Controllable construction of biocompatible supramolecular micelles and vesicles by water-soluble phosphate pillar[5]arenes for selective anti-cancer drug delivery. Chem. Mater. 28, 3778–3788 (2016).

43. Joseph, R. et al. Cationic pillararenes potently inhibit biofilm formation without affecting bacterial growth and viability. J. Am. Chem. Soc. 138, 754–757 (2016).
48. Pan, S. et al. Well-defined pillararene-based azobenzene liquid crystalline photoresponsive materials and their thin films with photomodulated surfaces. Adv. Funct. Mater. 25, 3571–3580 (2015).
49. Shi, B. et al. Nanoparticles with near-infrared emission enhanced by pillararene-based molecular recognition in water. J. Am. Chem. Soc. 138, 80–83 (2016).
50. Sun, Y., Guo, F., Zuo, T., Hua, J. & Diao, G. Stimulus-responsive light-harvesting complexes based on the pillararene-induced co-assembly of beta-carotene and chlorophyll. Nat. Commun. 7, 12042–12055 (2016).
51. Tan, L.-L. et al. Pillar[5]arene-based supramolecular organic frameworks for highly selective CO2-capture at ambient conditions. Adv. Mater. 26, 7027–7031 (2014).
52. Wang, S. et al. Thermo- and oxidation-responsive supramolecular vesicles constructed from self-assembled pillar[6]arene-ferrocene based amphiphilic supramolecular diblock copolymers. Polym. Chem. 8, 682–688 (2017).
53. Zhang, X. et al. Enzyme-regulated fast self-healing of a pillararene-based hydrogel. Biomacromolecules. 18, 1885–1892 (2017).
54. Luo, L. et al. Dynamic self-assembly adhesion of a paraquat droplet on a pillar[5]arene surface. Angew. Chem. Int. Ed. 55, 12713–12716 (2016).
55. Ni, M. et al. Dramatically promoted swelling of a hydrogel by pillar[6]arene-ferrocene complexation with multistimuli responsiveness. J. Am. Chem. Soc. 138, 6643–6649 (2016).
56. Jie, K., Zhou, Y., Yao, Y., Shi, B. & Huang, F. CO2-responsive pillar[5]arene-based molecular recognition in water: establishment and application in gas-controlled self-assembly and release. J. Am. Chem. Soc. 137, 10472–10475 (2015).
57. Ogoshi, T., Kida, K. & Yamagishi, T.-a Photoresponsive materials and their thin films by mesh printing. J. Mater. Chem. C 4, 8385–8391 (2016).
58. Liu, C. et al. Index-tunable anti-reflection coatings: Maximizing solar modulation ability for vanadium dioxide-based smart thermochromic glazing. J. Alloy. Compd. 731, 1197–1207 (2018).
59. Yan, Q. et al. Voltage-responsive vesicles based on orthogonal assembly of two homopolymers. J. Am. Chem. Soc. 132, 9268–9270 (2010).
60. Xia, W. et al. Responsive gel-like supramolecular network based on pillar[6]arene-ferrocene recognition motifs in polymeric matrix. Macromolecules 48, 4403–4409 (2015).
61. Xia, W., Hu, X.-Y., Chen, Y., Lin, C. & Wang, L. A novel redox-responsive pillar[6]arene-based inclusion complex with a ferrocenium guest. Chem. Commun. 49, 5085–5087 (2013).

Acknowledgements
This work was supported by the National Basic Research Program of China (2014CB846004) and the National Natural Science Foundation of China (No. 21472089 and 21572101). X.-Y.H. also thanks the Alexander von Humboldt Foundation for a research fellowship. We would like to thank Dr. Jochen Niemeyer form University of Duisburg-Essen for his very kind help on English language polishing.

Author contributions
L.W., X.-Y.H., and Y.-Z.S. conceived the project and supervised the research; S.W. designed and conducted the experiments; Z.X. performed parts of the synthetic experiments; T.W. performed the transmittance measurements; T.X. performed the drawing of the schematic diagram of hydrogels; S.W., X.-Y.H., and L.W. wrote the manuscript. All authors analyzed the data, discussed the results, and commented on the manuscript.

Additional information
Supplementary Information accompanies this paper at https://doi.org/10.1038/s41467-018-03827-3.

Competing interests: The authors declare no competing interests.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

© The Author(s) 2018