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Advanced liquid crystal-based switchable optical devices for light protection applications: principles and strategies

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Abstract

With the development of optical technologies, transparent materials that provide protection from light have received considerable attention from scholars. As important channels for external light, windows play a vital role in the regulation of light in buildings, vehicles, and aircrafts. There is a need for windows with switchable optical properties to prevent or attenuate damage or interference to the human eye and light-sensitive instruments by inappropriate optical radiation. In this context, liquid crystals (LCs), owing to their rich responsiveness and unique optical properties, have been considered among the best candidates for advanced light protection materials. In this review, we provide an overview of advances in research on LC-based methods for protection against light. First, we introduce the characteristics of different light sources and their protection requirements. Second, we introduce several classes of light modulation principles based on liquid crystal materials and demonstrate the feasibility of using them for light protection. In addition, we discuss the problems and shortcomings of current strategies. We propose several suggestions for the development of liquid crystal materials in the field of light protection.

Introduction

Human life is inextricably bound to light. Developments in science and technology have increased the popularity of artificial light and diversified the use of sunlight. Applications of artificial lighting are increasing at a rate of 6% per year^{1–3}. Simultaneously, laser technology has developed rapidly over the last three decades, and its range of applicability has expanded⁴. However, inappropriate light radiation can be harmful to human health and wellbeing^{5–12}. High-energy light can penetrate the lens of the human eye and reach the retina where it causes the retinal pigment epithelium to decline and thus affects vision. Additionally, for shorter wavelengths of visible light, the

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focal point does not fall at the center of the retina but rather slightly farther away. This creates a long-lasting state of tension within the eyeball, causing visual fatigue¹³. Furthermore, an increasing number of traffic accidents are caused by temporary "loss of vision" by drivers due to glare. In the United States alone, nearly 20 accidents occurred each day due to "laser dizziness" in 2017⁷. Considering that windows are the only means that allow outside light into a building or a vehicle, they play an important role in protection from light¹⁴. Consequently, it is important to design light protection for windows.

Throughout this review, the term "light protection" refers to means of preventing or weakening damage, discomfort or interference to people and photosensitive instruments by inappropriate light radiation. This can be accomplished by changing the intensity, energy, direction, and other characteristics of light passing through optical windows.

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Several types of light protection technologies have been investigated in recent years, including suspended particle devices (SPDs)^{15,16}, liquid crystal (LC) material systems^{17–25}, electrochromic devices (EDs)^{26–29}, photochromic devices 3^{30-32} and thermochromic devices 3^{3-37} . As shown in Fig. 1, each technology has different characteristics in terms of working principles, operating conditions, and performance. SPD films contain suspended needlelike or rod-like halide particles that align perpendicular to the substrate and allow light to pass through when an AC current is applied; they orient randomly and absorb or reflect light due to Brownian motion when unenergized. Thus, SPD films have fast switching speed but require electricity to maintain transparency¹⁶. Current EC materials can be divided into inorganic materials (functional transition metal oxides), organic materials (including small organic molecules and conductive polymers), and metal complexes. Their color change is achieved through redox reactions. When a suitable voltage is applied, an EC material captures injected electrons and is reduced and ionized, and its optical properties are changed. This color change consumes small amounts of energy, but the rate of color change is limited^{38,39}. When exposed to ultraviolet (UV) light, photochromic windows exhibit weak color switching for several minutes³¹. Most thermochromic devices are based on metal oxides, where the lattice structure of the material changes when the temperature reaches the phase transition temperature, and thus, the optical properties change. The high energy consumption of phase changing metal oxides as well as the limited visibility of thermochromic pigments restrict their widespread use⁴⁰.

As shown in Fig. 2, compared to conventional electronically controlled color-changing materials, LC materials exhibit superiority in terms of response time and protective bandwidth. Compared to conventional thermochromic materials, LC materials have great advantages in terms of stimulation temperature and visible light transmittance.



The LC phase is between crystalline solid and isotropic liquid states. This fascinating thermodynamically stable form of matter exhibits both the ordered nature of crystals and the fluidity of liquids. There are two types of LCs: thermotropic and lyotropic. LC phases form in thermotropic LCs because of temperature changes. Thermotropic LC phases are observed in rod-like, disc-like, and bent-core compounds. Rod-like thermotropic LCs are one of the most widely used materials in commercial applications ^{41,42}. The use of LCs in photo-regulation applications is assisted by their unique mix of reactivity to their environmental and stunning optical features, which allow immediate viewing of their response^{41,43}. Unless otherwise noted, throughout this paper, the term "LC" refers to rod-like thermotropic LCs.

The characteristics of light sources and their protection requirements in different light environments are discussed in the second part of this review. In the third part, principles of light protection using liquid crystal materials are described in detail. In the Section 'Strategies for LCbased light protection', light protection strategies based on liquid crystal materials that are suitable for individualized protection requirements in different light environments are discussed. Finally, opportunities for the development of light protection by liquid crystal materials are proposed.

Specific light environment confronted by LCbased light protection

External and internal light sources create an interior light environment. Natural and artificial light enter a room through windows. Light sources vary in frequency, energy, and irradiation area. They also pose different risks⁴⁴. Thus, it is necessary to discuss their characteristics to determine the best protection.

Protection against natural light

Natural light refers to light that is present in nature, such as sunlight, fire, and lightning flashes. Sunlight is the predominant source of natural light, and therefore, this section primarily discusses the light environment created by sunlight and the associated protection requirements.

As shown in Fig. 3a, the wavelength range of sunlight reaching Earth is from 250 to 2500 nm, which essentially covers from the UV to the near-infrared band. In sunlight, more than 90% of the energy is concentrated in the visible and infrared bands⁴⁵. Earth receives approximately 1373 W/m² of solar radiation, and the intensity of solar



radiation at its surface depends on the altitude and angle of the Sun, distance between the Sun and Earth, and duration of sunshine 46 .

Due to the rotation and revolution of the Earth, the duration of sunshine differs regionally⁴⁷. The radiation from the Sun on the ground plane consists of direct radiation and diffuse radiation. Direct radiation refers to radiation that comes directly from the Sun and does not change direction, whereas diffuse radiation refers to solar radiation that changes direction after being reflected and scattered by the atmosphere⁴⁴. Diffuse reflection changes not only the direction but also the intensity and spectral distribution of radiation. Solar irradiance varies sinusoidally throughout the day and fluctuates irregularly under the influence of clouds⁴⁸. In summary, the irradiation of sunlight has the characteristics of wide spectral range, large variation in irradiation angle, and wide area of irradiation.

The absorption of sunlight provides significant amounts of energy used by industries and daily life; however, it also has negative effects. Due to the conversion of heat by photoenergy, light exposure can cause an increase in the temperature of biological tissues. At increase of at least $10 \,^{\circ}$ C in temperature induces denaturation of many proteins in the retina, which results in photothermal damage⁴⁹. In addition, certain specific wavelengths of light radiation can cause chemical reactions in organisms, and photochemical damage can occur. Photochemical damage occurs when the retina is exposed to incident radiation in the high-energy portion of the visible spectrum⁵⁰. Several recent studies showed that photochemical damage occurred when the retina was exposed to incident radiation at high-energy wavelengths in the visible spectrum, which resulted in impaired vision and potentially serious ocular disease^{13,50}. At high latitudes or at sunrise and sunset, the altitude angle of the sun decreases, and consequently, it becomes easier for sunlight to enter the human eye obliquely, thus increasing the risks of accidents⁵¹. In addition, due to the wide irradiation area and various diffuse reflections, strong sunlight can affect the normal operation of light protection devices. That is, sunlight produces a large amount of noise and overwhelms the measurement signal, leading to device failure^{52–56}.

The wide distribution of sunlight necessitates special protection. Because insolation intensity varies over time, protective material must be adjustable⁵⁷. The changing angle of incident sunlight throughout the day necessitates

wide angle protective materials. Additionally, simultaneous protection from visible light and infrared radiation is required with broadband shielding over the 390–2500 nm range.

Additionally, protective materials must function throughout the day. The accumulation of solar energy raises the temperature of protective materials. This poses a serious threat to the properties of the materials. Additionally, for use during many hours of operation, protective materials must be energy efficient.

Protection from artificial light

Artificial light sources have aided human scientific and technological advancements. With the widespread use of artificial light sources, their effects on human life and health have increased. Lasers and other high-intensity lights are common sources of concern.

Lasers have played a groundbreaking role in many scientific and technological fields^{58,59}. Laser light is produced by transitions of electrons from excited to ground states⁶⁰. Instruments have been developed using different laser mediums, which lead to different wavelength distributions. Lasers primarily include single-frequency and multifrequency lasers⁶¹, as shown in Fig. 4a.

However, potential threats resulting from the use of lasers have increasingly attracted attention. Liu et al.⁵ presented the pathological effects of lasers at different wavelengths. The power of a laser can be as great as millions of watts per second, which can severely affect human vision (cause blindness) and destroy optical sensors. As a result, the use of lasers in weapons is frequently reported. Aside from irreversible eye damage, certain types of laser illumination may create transitory visual effects, such as distraction, confusion, or discomfort, and create potentially dangerous conditions for pilots of airplanes or drivers of vehicles^{62,63}. According to statistics from the Federal Aviation Administration (FAA), pilots in the United States reported observing laser light 9,723 times in 2021, as shown in Fig. 4b. This was a 42% increase over the 6,852 complaints received in 2020⁶².

As shown in Fig. 4c, several researchers systematically studied and calculated the possible forms of harm to people caused by different types of lasers (wavelength, beam propagation, and power) at various distances. Based on the likelihood of causing injury, various organizations, such as the American National Standards Institute, the Center for Devices and Radiological Health and the International Electrotechnical Commission, have classified lasers into four hazard classes^{64–66}. These data show that in the range of visible wavelengths, green lasers cause the strongest interference in the visual field, while with increasing distance, laser damage to people mainly produces transient visual effects and causes distraction, i.e., the first three levels of laser hazards. The use of optical

protection from high power lasers that may even cause fires is beyond the scope of this paper and therefore is not discussed further. As the light spot expands, the interference to the visual field of a driver gradually increases, as shown in Fig. 4d.

Owing to the high power, narrow band, and fast response of lasers, researchers have proposed different methods of protection from different types of lasers⁶⁷. As laser protection theory has continuously improved, requirements for protective material systems have become more specific. First, the protective band must cover the preset laser band range. Second, the protective effect should be adequately strong to provide protection from high-energy lasers. Finally, the materials must respond quickly to manage exposure to bursts of laser light.

In addition, high power lasers can increase the temperature of protective materials, which is a crucial factor in their failure. High-intensity light sources, such as sky lanterns and spotlights, are increasingly used in daily life, which results in a series of problems⁶³. Currently, the technologies that are primarily used for high-intensity lighting are carbon arc lamps, enclosed arc lamps, and high-intensity discharge lamps. The lighting characteristics of these technologies are listed in Table 1.

The most common problem is dazzling by highintensity lighting, as shown in Fig. 4e. Dazzling refers to visual conditions that can reduce the visibility of objects owing to an inappropriate brightness distribution or extreme brightness contrast in space or time. Dazzling originates from several sources, and therefore, its wavelength covers a wide spectral range. In addition, the duration of dazzling is short, and the intensity is relatively high intensity, which causes discomfort to personnel in buildings or vehicles and results in severe consequences.

The primary requirement for protection against dazzling by light is fast response speed that is comparable to that of lasers. Additionally, high-intensity lighting has a wider spectral range than lasers, including the entire visible light band, and this requires protective equipment that has a wider range of spectral adjustment.

Currently, there are several types of light protection devices based on LC materials, and each has different effects. Smart optical devices using LC materials can realize multiband protection, light intensity regulation, and fast responses to multiple stimuli, among other functions. Consequently, LC materials⁶⁸ have been widely used in light protection⁶⁹. The next section discusses the application of LCs in the field of light protection from the perspective of principles of protection.

Principles of LC-based light protection

LC-based materials for light protection should correspond to the spectral distribution, intensity, and angle of



incidence for specific surroundings. In this section, LCbased materials are classified based on different light protection mechanisms. We focus primarily on the conversion and transfer path of photoenergy. In Section 'Absorption-based light protection', we introduce LCbased materials with light energy conversion as the working mechanism. These materials apply the principle of light absorption to provide protection. In Section 'Reflection-based light protection', we demonstrate photoenergy transfer by the directional path of LC-based materials for light protection. Here, the focus is on reflection by LC-based materials. In Section 'Scatteringbased light protection', we discuss protection by photoenergy transfer by nondirectional paths in LC-based materials based on the scattering characteristics of LCbased materials. Finally, we summarize the properties of LC-based materials based on different light protection mechanisms in complex environments.

Searchlight	Diameter searchlight (m)	llluminance at 100 m (lx)	Luminous intensity (cd)	Luminance (cd/m ²)	Beam width (deg)
Arclight	1.524	5200	5.2×10^{7}	2.87×10^{7}	0.93
Dominator	1.524	14000	1.4×10^{8}	7.72×10^{7}	2.93
Prolight	0.406	727	7.27 × 10 ⁶	5.60×10^{7}	1.68
Xenon	0.406	7013	7.01×10^{7}	5.41×10^{8}	1.72
Skytracker	0.406	3320	3.32×10^{7}	2.56 × 10 ⁸	2.67
Carbon arc	1.524	12900	1.29 × 10 ⁸	4.11×10^{9}	2.85

Table 1 Lighting characteristics/parameters of various high-intensity lights/technologies⁶³



Absorption-based light protection

The active form of photoenergy conversion involves converting photoenergy into heat energy, chemical energy, and/or other forms of energy through absorption by LC-based materials. As shown in Fig. 5a, the absorption bands of the thermally induced rod-like LC molecules are concentrated in ultraviolet (<380 nm) and long wavelength infrared light (>3 μ m)⁷⁰. In recent years, good light protection was obtained by mixing functional

materials with visible and near-infrared absorption properties and LCs.

Electron transition-based absorption

Each molecule has its own spectral transitions. Light radiation is absorbed by a molecule when it resonates with electronic transitions, as shown in Fig. 5b. The transitions of valence electrons in several organic molecules can absorb light in the wavelength range of 200–1000 nm,

which lies in the UV–visible region⁷¹. Therefore, the most straightforward approach for obtaining light absorption is to dope the host LC with organic dyes.

Light absorption by a dye molecule is directional or dichroic. There are two types of dichroic dyes, those with positive and negative absorption characteristics. Positive (p-type) dichroic molecules absorb polarized light best along the long axis and almost no light polarized perpendicular to the long axis, as shown in Fig. 5c. Negative (n-type) dichroic dyes have the opposite behavior⁷².

Meanwhile, dichroic dye molecules have rod-like shapes that are similar to those of LC molecules. When dichroic dyes are dissolved in an LC host, the LCs act not only as solvents but also as agents of structural alignment. By adjusting the LCs, the orientation of the dye molecules can be changed, as shown in Fig. 5d; this is known as the guest-host effect. Since the first LC display based on the guest-host effect was reported by Heilmeier et al. in 1968, this type of device attracted widespread attention and was gradually incorporated into light protection devices by researchers⁷³.

While many dichroic dyes were developed, none were synthesized with uniform absorption across the entire visible light spectrum, which limited the use of the guest-host effect of LC materials in light protection applications. Thus, multiple dichroic dyes were mixed to formulate guest-host LCs with broad absorption bands, as shown in Fig. 6a, and thereby achieve absorption across almost the entire visible range of the spectrum^{74–77}. In addition, the solubility of dichroic dyes in LCs and the order parameter (S) greatly affect the performance of guest-host light protection devices. Both factors determine the absorbance and contrast of the device^{78–85}, as shown in Fig. 6b. However, this type of research primarily involves the structural design of dye molecules, which is not discussed in this paper.

Additionally, dye molecules absorb only polarized light, which makes them inefficient light absorbers. Light is strongly absorbed when its polarization is parallel to the long axis of a dye molecule. When light is oriented perpendicular to the long axis of a dye molecule, it is weakly absorbed⁸². One method to improve the absorption efficiency is to design a double-layer structure wherein the LC molecules of the upper and lower layers are arranged perpendicular to each other, as shown in Fig. 6c. LC molecules in this arrangement absorb light with all polarizations⁸⁶. Another method is to change the alignment of liquid crystal molecules. Huh⁸⁷ et al. investigated the angular dependence of dye transmittance in planar Ch-LC, PSLC, and ion-doped LCs, as shown in Fig. 6d. A twisted structure in the planar state of the Ch-LC cell resulted in the least transmittance. Further studies found that considering the wave-guide effect, the transmission in the planar state decreases as the number of pitches in the Ch-LC cells increased for the same cell gap⁸⁸, as shown in Fig. 6e. However, this required the addition of more chiral dopants, which led to an increase in the driving voltage.

Surface plasmon resonance-based absorption

As shown in Fig. 7a, nanoparticles (NPs) exhibit surface plasmon resonance at specific wavelengths with strong absorption effects because an evanescent wave forms in the optically sparse medium and a specific plasma wave forms in the medium when light is totally reflected by the surfaces of the NPs. When two waves meet, resonance occurs. When the evanescent wave resonates with the surface plasmon wave, photons gain energy. The surface plasmon wave absorbs most of the incident light energy, which reduces the reflected light energy^{89–91}. The idea of doping functional particles has been a hot topic in liquid crystal research since 1970, when F. Brochard and PG de Gennes modified the properties of liquid crystals by mixing them with submicron magnetic particles⁹². In recent years, with the continuous development of nanotechnology, liquid crystal light protection devices doped with nanoparticles have attracted attention.

The effect of nanoparticles on LCs were found to depend on several factors, such as material composition, shape, size, concentration, modification form and liquid crystal type⁹³. Inorganic semiconductor NPs displayed excellent UV and infrared shielding properties due to their localized surface plasmon resonance (LSPR). This suggested new possibilities for plasmonic light manipulation by altering the material composition of doped semiconductors^{94–96}. In addition, anisotropic gold nanorods (GNRs), important nanostructures with remarkable optical properties, were investigated⁹⁷. The surface plasmon resonance of anisotropic GNRs enabled them to concentrate and manipulate light based on their size, shape, and proximity. Similar to the guest-host effect described above, thiol molecules were introduced on the surface of GNRs so that they could be arranged with LCs, as shown in Fig. 7b⁹⁸⁻¹⁰¹. Electrical switching of both visible and infrared light transmission using an LC film containing a dichroic dye and GNRs was demonstrated^{102,103}. As shown in Fig. 7b, doped dye molecules and GNRs spontaneously aligned along or across the director field of the host LC. The composite film absorbed light from visible to near-infrared and switched quickly at low voltage.

However, the LC/monomer mixture is usually viewed as a solvent with high polarity. This drastic change in the polarity of the medium destabilizes the dispersion and leads to severe aggregation. In studies, this not only resulted in the loss of transmittance in the visible region but also detrimentally affected the near-infrared shielding ability of NPs owing to the significant reduction in the surface electron density in large aggregates^{104–106}. As shown in Fig. 7c, modification of NPs by functionalization or addition of solvents prevented aggregation effects⁹³. For example, ITO



NPs were encapsulated with an insulating silica barrier to form a core/shell structure to prevent them from aggregating²¹ (Fig. 7d). However, this approach had an impact on the absorption spectra of NPs due to the surface chemistrydependent modulation of the LSPR of NPs¹⁰⁷. Additionally, the dispersion of NPs in LCs significantly affected the various physical and electrical properties of the host LC material¹⁰⁸. Therefore, for devices that rely on the electrooptical response of liquid crystal molecules, it is necessary to consider the multiple effects of NP doping.

The advantage of the photoenergy conversion method of LC-based light protection materials is that they can completely consume light by converting photoenergy into other forms of energy. Therefore, this method provides a good light protection effect; however, new forms of energy may cause a series of problems and failure of the material.

Reflection-based light protection

Cholesteryl liquid crystals (Ch-LCs), the most typical reflective material, were discovered in 1888 in the form of cholesteryl esters^{109,110}. They spontaneously exhibit helical superstructure and can selectively reflect up to 50% of

unpolarized natural light. They are generally developed by introducing chiral molecules into achiral nematic LCs. Chiral dopants generate LC organization, wherein successive layers of nematic LCs are displaced by a small rotation in the molecular director with respect to neighboring layers. The resulting "twist" may be either right- or left-handed. In the cholesteric phase, the molecules possess a helical distribution with a periodic helical structure. According to Bragg's law shown in Fig. 8a, the central reflection band of Ch-LC is determined by the pitch (P), as shown in Fig. 8b, average refractive index (n_{avg}) of the material, and incident angle of the light. The pitch is defined as the distance over which the director field rotates by one full turn (360°). The pitch (P) of the Ch-LC depends on the concentration (C) and helical twisting power (HTP) of the chiral dopants (Eq. 1).

$$P = \frac{1}{C \times \text{HTP}} \tag{1}$$

The bandwidth of the light reflected by the Ch-LC is determined by the difference between the extraordinary



 (n_e) and ordinary (n_o) refractive indices as well as the pitch of the host LC (Eq. 2), which is normally limited to between approximately 75 and 100 nm¹¹¹.

$$\Delta \lambda = (n_e - n_o) \times P \tag{2}$$

The maximum reflection produced by the cholesteric reflector layer is limited to 50% of the unpolarized natural light because Ch-LCs can only reflect circularly polarized light whose polarization matches that of the helix. For example, right-handed cholesteric light reflects only right-circularly polarized light, as shown in Fig. 8c. Both left-circularly polarized light and light outside the cholesteric reflection bandwidth are unaffected by the LC matrix and are transmitted normally¹¹². It is necessary to overcome these limitations in the fabrication of polarization-independent devices, such as light protection devices¹⁰⁵.

To broaden the bandwidth around the central reflected wavelength, the most straightforward approach is to stack single Ch-LC layers with different pitch lengths. Both Hideo Takezoe et al. and Shin-Tson Wu et al. constructed multi-CLC systems and introduced phase-modulated materials to overcome the reflection limit^{113,114}. In addition to this multilayer stacking approach, Mitov¹¹⁵⁻¹¹⁸ et al. used polymer-based composites to construct pitch gradients in a single Ch-LC layer, as shown in Fig. 8d. They coated two glass plates with cholesteric films of different pitches and used thermal diffusion between the layers to create a lateral concentration gradient of the chiral agent in the LC unit, which resulted in a gradient of spacing in the Ch-LC layer and an expanded reflection band. In addition, several studies used UV-induced reactive polymer monomers to generate UV intensity gradients in films by adding UV-absorbing dyes to blends of



raw materials to alter the polymerization rate within the films and thus construct spacing gradients in the Ch-LC layers^{119–125}. Zhou¹²⁶ et al. added a DC bias during the photopolymerization process. This bandwidth broadening was attributed to a frozen pitch gradient that resulted from an in situ electric-field-assisted dynamic ion-dragging effect, which led to the formation of a pitch gradient along the direction of the electrical field.

To break the reflection limit, Guo¹²⁷ et al. employed a "wash-out/refill" technique to use polymers to stabilize LC films with two-handed circularly polarized reflection bands, as shown in Fig. 8e. In contrast, Mitov¹¹⁷ et al. used thermally induced helicity-inverted Ch-LCs to prepare LC films stabilized with polymers; the maximum reflectivity of the films was 72%.

In conclusion, by using polymer-liquid crystal (P-LC) materials, the inherent defects of Ch-LC can be greatly improved, and the application of this approach in the field of light protection can be expanded.

According to previous research, photoenergy transfer by the reflected path of LC-based materials adjusted the propagation path of light through reflection to provide light protection. This method had no secondary negative consequences of energy conversion, and the light attenuation rate was high. However, the reflective property depended on the structure of the material, had inherent angular dependence and required films with high dimensional accuracy.

Scattering-based light protection

As shown in Fig. 9a, propagation of light in materials with refractive index inhomogeneities results in scattering. Multiple scattering events cause photoenergy to deviate from its original propagation direction and spread out for transmission. As shown in Fig. 9b, LCs with controllable birefringence may provide an effective means for adjusting the refractive index of the medium. Depending on the refractive index adjustment method, protective materials can be divided into two categories: static and dynamic scattering materials.

Static scattering

Polymer-liquid crystal (P-LC) materials were used for fabricating commonly commercialized haze-switchable LC smart windows¹²⁸. The P-LC protective system was divided into polymer-dispersed liquid crystal (PDLC) and polymer-stabilized liquid crystal (PSLC) systems depending on the polymer content in the system¹²⁸. In the PDLC system with high polymer content, the LC was dispersed within the polymer matrix in the form of small microdroplets. The LC molecules within the micro-droplets were freely oriented. Their refractive indices did not match the refractive indices of the matrix and presented a scattering state. The direction of the optical axis of the liquid crystal microdroplet was adjusted by applying an electric field. When the refractive indices of both matched, a transparent state formed, as shown in Fig. 9c.



In the 1980s, J. W. Doane et al. first prepared PDLC films by a phase separation method¹²⁹. This technique was extensively utilized in both academic research and industrial development of polymer-liquid crystal (P-LC) materials. For PSLC systems with low polymer contents, the polymer bound some of the LC molecules, and its optical axis direction was different from that of the LC molecules in other regions, which resulted in a refractive index mismatch and a scattering state¹³⁰. In addition, cholesteric LCs exhibited metastable scattering textures and were transformed into a focal conical domain by external stimuli, as shown in Fig. 9d^{130,131}. Therefore, Ch-LCs are often used in P-LC systems to enhance this effect.

Optimizing the performance of LC/polymer materials has been the focus of research in recent years. Several key aspects used to evaluate the electro-optical performance of LC/polymer materials were the driving performance (threshold voltage V_{th} and saturation voltage V_{sat}), the response time (rise time τ_{on} and decay time τ_{off}) and the optical contrast (ratio of transmittance in the on-state and off-state of the film). Theoretically, the driving performance of PDLC films and the response time can be predicted according to Eqs. $3-6^{132-136}$.

$$V_{\rm th} \approx \frac{d}{R} \times \left[\frac{K(\omega^2 - 1)}{\varepsilon_0 \Delta \varepsilon} \right]^{\frac{1}{2}}$$
(3)

$$V_{\rm sat} \approx \frac{d}{R} \times \left(\omega^2 - 1\right)^{\frac{1}{2}} \times \frac{4\pi K}{\Delta \varepsilon}$$
 (4)

$$\tau_{\rm on} \approx \frac{\gamma}{\Delta \varepsilon V^2 - \frac{K(l^2 - 1)}{R^2}} \tag{5}$$

$$\tau_{\rm off} \approx \frac{R^2 \gamma}{K(l^2 - 1)} \tag{6}$$

where *V* is the applied electric field, while *d*, *R*, *K*, ω , ε_0 , $\Delta \varepsilon_1$ *l*, and *y* represent the film gap, droplet radius of the LC, elastic constant, aspect ratio, vacuum dielectric constant, dielectric anisotropy, shape anisotropy, and rotational viscosity constant, respectively. The addition of NPs changed the refractive index of the dielectric, the dielectric constant, and the anchoring properties of the liquid crystal, and thus, effectively improved the electro-optical properties of LC/polymer materials^{130,137}. For example, SO_2 NPs changed the anchoring energy at the LC-polymer interface, and inorganic NPs such as ZnO and MgO changed the dielectric constant of the dielectric, both of which effectively reduced the driving voltage. Due to the surface plasmon excitation at the metal-LC interface, which enhanced the local electric field, films doped with metal NPs had low driving voltages and high CR values^{130,138–140}. However, as mentioned above, the dispersion stability of NPs in liquid crystals is still a major challenge. In addition to the inherent properties of the liquid crystal molecules,



the polymer morphology affects these properties. For example, Li et al.¹³⁶ modified polymer microstructure and altered the electro-optical properties of films by adjusting the content of epoxy resin monomer, as shown in Fig. 10a. Reducing the LC droplet size shortened the response time but it also increased the driving voltage.

For "normal mode" P-LC films, the films are opaque in the absence of an applied field (off-state) and become transparent in the presence of an applied field (on-state). Yao-Dong Ma¹⁴¹ et al. proposed the opposite operation of P-LC films, called the reverse mode, where films were transparent in the off-state and changed to opaque in the ON state when a small electric field was applied. This type of film helped to broaden the range of applications of P-LC materials and attracted the attention of many scholars. In recent years, Meng et al. proposed a reverse-mode PSLC (R-PSLC) smart window that used a substrate with an uneven surface, as shown in Fig. 10b; it significantly increased the haze by 42% compared to that for a substrate with a smooth surface¹⁴². Deng et al. prepared a trans-PDLC smart window using a carbon nanotube as the alignment layer¹⁴³. The optical performance of these smart windows was adjusted using near-infrared irradiation to fabricate highly efficient responsive PSLC smart windows without additional materials for orientation. This improved the durability of the device.

Thermally responsive P-LC materials were prepared based on the liquid crystal phase transition. Hiroshi Kakiuchida¹⁴⁴ et al. developed a nonuniform irradiation method for photopolymerization-induced phase separation. The transparent and scattering states of LC/polymer films were controlled by temperature-induced phase separation of nuclei, and thermally responsive polymer network liquid crystal films in normal mode and reverse mode were developed, as shown in Fig. 10c.

Dynamic scattering

As early as the late 1960s, researchers found that when low-frequency electric fields were applied to some



nematic liquid crystals, the LCs exhibited significant hydrodynamic instabilities. When the voltage exceeded a threshold value, as shown in (Fig. 9e), turbulence developed within the liquid crystal layer and produced strong scattering of light^{145,146}. One of the main problems with the electrical characteristics of the dynamic light scattering mode was the high operating voltage;¹⁴⁷ the predicted threshold voltage is given by Eq. 7¹⁴⁸,

$$V_0^2 = \frac{\pi^2 (1 + \omega^2 r^2) k_{33} \sigma_{\parallel}^2 q^2}{\varepsilon_0^3 \varepsilon_{\parallel} \varepsilon_{\perp} \varepsilon_{\alpha} (\omega^2 - C)}$$
(7)

where

$$C = \frac{\sigma_{\parallel}}{\varepsilon_0^2 \varepsilon_{\perp}} \left[\frac{-\alpha_2}{\eta_1} \left(\frac{\sigma_{\parallel}}{\varepsilon_{\parallel}} - \frac{\sigma_{\alpha}}{\varepsilon_{\alpha}} \right) - \frac{\sigma_{\perp}}{\varepsilon_{\parallel}} \right]$$
(8)

ω is the angular frequency of the applied voltage, τ is the charge relaxation, k_{33} is the bend elastic constant, q is the wavenumber (approximately π/d), $σ_{\perp}$ and $σ_{\parallel}$ are the

perpendicular and parallel components of the conductivity, σ_a is the conductivity anisotropy ($\sigma_{\alpha} = \sigma_{\parallel} - \sigma_{\perp}$), ε_{\perp} and ε_{\parallel} are the perpendicular and parallel components of the dielectric constants, ε_{α} is the dielectric anisotropy ($\varepsilon_{\alpha} = \varepsilon_{\parallel} - \varepsilon_{\perp}$), and η 1and α_2 are shear viscosities.

Increasing the overall conductivity by adding dopants is an effective strategy to lower the threshold value. Zhan¹⁴⁹ et al. doped different ions in one LC material and discovered that the positive charge of the organic part in the salt was the primary cause of turbulence in the LC. Furthermore, Zhan et al. fabricated light-induced EHDI devices by doping LCs with a photochromic dye, spiropyran, as shown in Fig. 11b. With UV irradiation, the closed spiropyran ring opened to form open cyanocyanine (365 nm, purple flash). Ring closure was achieved with visible light (565 nm, green light). Tens of seconds of exposure to UV and green light activated or eliminated the EHDI effect¹⁵⁰. However, this technology was limited by long-term stability, heating, and high power consumption due to the accumulation of ionic charges at the interface between the LC and alignment layers or the movement of ions in the cell^{151,152}. To address these issues, Yoon¹⁵³ et al. proposed an LC smart window device based on the EHDI effect without ion doping. As shown in Fig. 11c, N-LC was used as the host material, and SmC*-LC was used as the guest material. The dielectric properties of the two materials were opposite, and incident light was strongly scattered by randomly oriented LC molecules when an electric field was applied, which led to a very hazy translucent state.

The scattering process can adjust the propagation path of light to produce light protection. The advantage of this method is that it can provide a wide viewing angle and yet it exhibits the protective effect for wideband light sources. A disadvantage is that the attenuation rate of light is low, and most light passes through the material surface via a scattering path.

These three types of traditional light protection mechanisms have their own characteristics and advantages. Devices prepared by these mechanisms were utilized to provide protection from different light sources in the past few decades and achieved good effects. The properties of LC materials based on different light protection mechanisms are summarized in Table 2.

However, these traditional mechanisms for light protection have inherent defects. With the increasing complexity of the light environment, we need to make comprehensive use of these mechanisms and combine their advantages to meet more demanding application requirements.

Strategies for LC-based light protection

Thus far, we have covered the characteristics of various light sources and LC-based light protection mechanisms. However, the requirements for light protection vary according to the light environment. In most cases, comprehensive protection requires multiple coupled mechanisms to meet requirements, such as fast response, high contrast, large-area fabrication, and energy efficiency. In various light protection scenarios, LC-based light protection strategies are different. As shown in Fig. 12a,b, transient light protection is often needed for vehicles. Buildings require devices that provide long-term light protection (Fig. 12c).

Transient light protection strategies

Traffic injuries are a significant cause of death in the world¹⁵⁴. The most common situations occur when driving while facing light, as when sunlight shines directly through the windshield into the eyes of the driver (pilot) and affects their vision. The conventional protection strategy is to use sun visors. However, as shown in Section 'Protection against natural light' above, the irradiation angle of sunlight varies greatly, and the use of visors in

fixed positions greatly limits their protective effect and is a safety hazard that could contribute to collisions. A better protection strategy is to use switchable light-shielding materials in local areas of the windshield to change the propagation path of incident light energy and reduce the damage caused by direct sunlight entering the eyes of drivers (pilots), as shown in Fig. 12a. Additionally, it is necessary for the windshield, as the main channel through which the driver (pilot) sees the outside world, to have a good visual field (high transmittance, low haze) and fast switching speed of light protection.

In Section 'Scattering-based light protection', P-LC materials based on the scattering principle exhibited protection from wideband light sources, which was consistent with the characteristics of the broad-spectrum radiation of sunlight. Therefore, these materials are potential candidates for windshield light protection. However, the reported switching speeds of commercial P-LC materials were typically within hundreds of milliseconds^{128,130,131,136,142–144,155–162}, and for important windows such as windshields, faster switching speeds are often sought.

The new protocol based on the doping of LC with a low concentration of nanoporous microparticles presented by the group of I. Abdulhalim created windows that rapidly switched between transparent and strong scattering states. In the OFF state, the average refractive index of these particles was highly mismatched with the refractive index of the surrounding LC region due to the random orientations of the particles, which produced strong scattering. In the ON state, the molecules oriented in the direction of the electric field, which increased the transparency of the system. As shown in Fig. 13a, researchers used three types of nanoscale and microscale particles to demonstrate this concept: (1) cochleate particles¹⁶³, (2) colloids of decanol¹⁶⁴, and (3) nanoporous Si microparticles¹⁶⁵. In the off and on states, the rise and fall times of these windows were approximately a few milliseconds and tens of milliseconds, respectively. For security reasons, a windshield should be completely transparent during an emergency. However, all the aforementioned methods were opaque at zero electric field; thus, they are unsuitable for this application.

Chun-Wei Chen and colleagues reported the development of a light protection window based on cholesteric LCs with negative dielectric anisotropy, as shown in Fig. 13b. The transparent state exhibited well-aligned planar cholesteric texture and was stable in the absence of an electric field, whereas the scattering state turned on when a field above the fluctuating instability threshold was applied. The material had 86% transmittance in the transparent state and could switch to the scattering state within 10 to 100 ms, where the specular transmittance decreased to 11%¹⁶⁶.

Mechanism	Stimulation (range, amplitude, action time)	Light modulation type	Optical modulation effect (range, action effect)	Recovery	Full recovery	Ref
Absorption-based light protection	ES ^a (AC ^b , 34 V, 100 ms)	Amplitude adjustment	800—2500 nm (S% ⁱ > 9 5%), 380–780 nm (T% ⁱ 83%—2%)	V ^I Off	\checkmark	176
	ES ^a (AC ^b , 30 V, 100 ms)	Amplitude adjustment	800–5500 nm (∆ T% ^j 20%), 380–780 nm (T% ^j 20%–78.5%)	V ^I Off, 300 ms	\checkmark	177
	TS ^d (~42 °C)	Amplitude adjustment	380–780 nm (T% ^j 80%–1%)	RT ^m	\checkmark	181
	TS ^d (27~35 °C)	Amplitude adjustment	380–780 nm (T% ^j 40%–0%)	RT ^m	\checkmark	182
	TS ^d (20~50 °C)	Amplitude adjustment	380–780 nm (T% ⁱ 83.6%—0) & Infrared band (Δ T% ^j 20%)	20 °C	\checkmark	179
Reflection -based	TS ^d (25∼75 °C)	Band adjustment	$2 \times RBs^k$	RT ^m	\checkmark	195
light protection	TS ^d (30~75 °C)	Band adjustment	RB: ^k 450 nm—660 nm	RT ^m , 3 h	\checkmark	196
	TS ^d (25~48 °C, 120 s)	Band adjustment	RB: ^k 440 nm—660 nm	-	-	171
	TS ^d (30~75 °C)	Band adjustment	RB: ^k 450 nm—640 nm RB: ^k 575 nm-650 nm	RT ^m , 3 h	\checkmark	196
	TS ^d (17~130 °C)	Band adjustment	RB: ^k 940 nm—1250 nm	RT ^m	×	196
	LS ^e (365 nm, 120 s)	Band adjustment	RB: ^k 440 nm—660 nm	LS ^e (450 nm)	\checkmark	171
	LS ^e (530nm, 60s)	Polarization adjustment	Reflected \rightarrow Transparent	LS ^e (440 nm)	×	197
	LS ^e (530 nm, 60 s)	Polarization adjustment	Reflected → Transparent	LS ^e (530 nm)	×	197
	LS ^e (360 nm, 2~10 min)	Band adjustment	RB: ^k 400–570 nm	Light Off, 24 h	\checkmark	169
	LS ^e (365 nm, 7~43 s)	Band adjustment	RB: ^k 400 nm—740 nm	LS ^e (520 nm),12 s	\checkmark	170
	LS ^e (365nm, 60s)	Band & Polarization adjustment	RB: ^k 300~420 nm −420~550 nm Right-handed-circularly → Left-handed-circularly	LS ^e (380–780)/Light Off/TS ^d	\checkmark	198
	LS ^e (980 nm, 120s)	Band adjustment	RB: ^k 400 nm—740 nm	LS ^e (980 nm), 120 s	\checkmark	199
	ES^{a} (AC ^b , 8.6 V μm^{-1})	Polarization adjustment	700–1400 nm Reflected → Transparent	V Off	\checkmark	19
	ES ^a (DC ^c , 0~1.2 V μm ⁻¹)	Band adjustment	RB: ^k 120 nm—1100 nm	V Off	\checkmark	200
	${\rm HS}^{\rm f}$ (RH ^g > 45%)	Band adjustment	RB: ^k 1150 nm—1570 nm	TS ^d (>30 °C, 5 min)	\checkmark	201
	PHS ^h (PH = 7~9, 3~5 min)	Band adjustment	RB: ^k 450 nm—550 nm	-	-	202
	PHS ^h (acidic, >200 s)	Band adjustment	RB: ^k 600 nm—770 nm	-	-	203
	PHS^{h} (PH = 9)	Band adjustment	RB: ^k 520 nm—720 nm	PHS ^h (PH 3)	×	204
	TS ^d (>60 °C)	Band adjustment	RB: ^k 648 nm—460 nm	RT ^m , 3 days	\checkmark	205
	TS ^d (>75 °C)	Band adjustment	RB: ^k 780 nm—540 nm	RT ^m	\checkmark	195
Scattering-based light protection	ES ^a (AC ^b , 100 Hz, 40 V)	Amplitude adjustment	T% ^j 64%—10%	ES ^a (AC ^b ,1000 Hz,15 V)	\checkmark	180
	ES ^a (AC ^b , ~20v, 84 ms)	Amplitude adjustment	T% ^j 60.8%—1.3%	V ^I Off	\checkmark	175
	ES ^a (AC ^b , 0.85~1.45 V/μm)	Amplitude adjustment	T% ^j 0.41%—98.98%	V ^I Off	\checkmark	167
	ES ^a (AC ^b ,1 kHz, ~3.8 V/ μm, 10~100 ms)	Amplitude adjustment	T% ^j 90%—20%	V ^I Off ES ^a (AC ^b ,1000 Hz, <2.8 V/μm)	\checkmark	166
	ESª (AC ^b , 55 V, 55 ms)	Amplitude adjustment	T% ^j 79%—1%	V ^I Off, 55 ms	\checkmark	184

Table 2 Categories of light protection mechanisms based on LCs

Mechanism	Stimulation (range, amplitude, action time)	Light modulation type	Optical modulation effect (range, action effect)	Recovery	Full recovery	Ref
	ES ^a (AC ^b , 24 V, <5 ms)	Amplitude adjustment	400 nm—800 nm (T% ^j ~0% —70%)	V ^I Off, 1000 ms	\checkmark	206
	ES ^a (AC ^b , 70 V, 40 ms)	Amplitude adjustment	Scattering \rightarrow Transparent	V ^I Off, 200 ms	\checkmark	164
	ES ^a (AC ^b , 40 V)	Amplitude adjustment	Scattering \rightarrow Transparent	V^{I} Off	-	207
	LS ^e (UV, 2 mW·cm ^{−2} ,30 s)	Amplitude adjustment	T% ^j 79%–1%	Light Off	\checkmark	180
	TS ^d (37.8 °C)	Amplitude adjustment	T% ^j 64%-16.7%	TS ^d (25°C)	\checkmark	184
	TS ^d (>30 °C)	Amplitude adjustment	380–780 nm (∆T% ^j 10%)	RT ^m	\checkmark	178
	LS ^e (980 nm, ~10 s)	Amplitude adjustment	Transparent \rightarrow Scattering	Light Off, ~30 s	\checkmark	25
	LS ^e (UV, 30 s)	Amplitude adjustment	Scattering \rightarrow Transparent	Light Off	\checkmark	207
	TS ^d (29 °C)	Amplitude adjustment	T% ^j 64%–17.1%	TS ^d (21.2 °C)	\checkmark	180

Table 2 continued

^aES Electrical Stimulation, ^bAC Alternating Current, ^cDC Direct Current, ^dTS Temperature stimulation, ^eLS Light stimulation, ^fHS Humidity stimulation, ^gRH Relative humidity, ^hPHS Potential of hydrogen stimulation, ⁱS% Shielding rate, ^jT% Transmittance, ^kRB Reflection band, ^lV Voltage, ^mRT Room temperature

Scattering-based equipment causes radiation to be scattered in the forward direction^{163,164,166}, and protection is not effective under strong sunlight. The light protection method of multi-mechanism integration combines traditional light energy transformation, reflected transfer, and scattering transfer, and it was widely applied in recent years.

Many scholars recently conducted research on doping dye molecules into P-LC materials to combine traditional light energy transformation and scattering transfer mechanisms to enhance the protection of the materials. Anuja Katariya-Jain¹⁶⁷ et al. investigated the effects of doping LC materials with dyes on the properties of the resulting PDLC films, as shown in Fig. 13c; the dopants changed the specular transmittance of the films from 0.41% to 98.98% at a low voltage (1.45 V/µm). A dye-doped R-PSLC smart window with a special electrode pattern was designed by Tae-Hoon Yoon¹⁶⁸ et al. As shown in Fig. 13d, the visible transmittance of this device changed from 67.0% (in the OFF state) to 3.2% (in the ON state) with a fast response time (<1 ms) and low operating voltage (10 V).

Artificial light sources, such as lasers and other forms of high-intensity lighting, also pose risks. As shown in V in Fig. 12a, traditional laser protection strategies are achieved by selectively blocking specific wavelengths of laser light. However, this causes difficulty for the driver (pilot) in distinguishing certain colors and affects visual performance. Additionally, since the glare interference source is uncontrollable, the material should react automatically with a passive protection strategy so that it is not necessary for the driver (pilot) to actively adjust the optical properties of the windshield. Currently, there are no proven protection technologies that provide this capability. However, one potential method involves using lightdriven chiral molecular switches to induce Ch-LCs to achieve reversible selectivity over the entire visible spectral range, a method known as reflective color tuning (shown in Fig. 14a)^{169,170}. The HTP value of the chiral molecules changes dramatically after photoisomerization. To achieve light protection when the device is exposed to a certain wavelength band (450 nm), the center of the reflection wavelength of the device shifts to 440 nm¹⁷¹.

In addition to windshields, optical windows on vehicles include aircraft portholes, vehicle side windows, sunroofs and side windows on ships. Unlike windshields, these windows primarily enhance the passenger experience and do not require windshield-level visibility and rapid switching speed for light protection. Due to the need for passengers to read and rest, it is necessary for these windows to have good thermal and light insulation properties, as shown in Fig. 12b. Because these windows may be in an opaque state for a long time, low energy consumption is also an issue that should be considered.

As mentioned above, dye-doped P-LC materials were capable of both light scattering and absorption and had good dimming properties. However, there was a trade-off between light scattering and absorption. Tae-Hoon Yoon⁸⁶ et al. used a negative LC with dye, and electro-hydrodynamic effects to provide an LC cell with a very low overall transmittance, as shown in Fig. 15a. At 40 V, the specular transmittance changed from 60.1% to 3% with strong scattering. In addition, in dye-doped P-LC materials, the dye dissolved in the polymer matrix, which limited the contrast. Kim et al. reported encapsulating dye in monodisperse capsules to solve dye-related issues. A contrast ratio of 120 or greater was achieved after encapsulation, as shown in Fig. 15b¹⁷². When preparing



dye-doped PSCLC films, some studies combined traditional light energy conversion, reflected transfer, and scattering transfer¹⁷³. The device shown in Fig. 15c switched between reflected green light, black fog, and transparent states with a response time of hundreds of milliseconds.

Unlike visible wavelengths, light transmission in the infrared wavelengths brings heat radiation, so it is essential to manufacture windows that possess IR shielding properties along with good transparency. A hybrid window made of nanoscale and microscale LCs and a layer of thermochromic material (VO₂) was

demonstrated recently to fulfill this purpose, as shown in Fig. 16a¹⁷⁴. Furthermore, many researchers tried doping NPs into P-LC materials to expand the modulation band while ensuring the fast response of light protection devices. Liang¹⁷⁵ et al. reported the development of an LC light protection device with controllable visible and near-infrared light transmittance, which contained microscale LC domains with a negative dielectric constant and tungsten-doped vanadium dioxide (W-VO₂) nanocrystals. Due to the combination of scattering with absorption and reflection, the transmittance of near-infrared light was reversibly and passively modulated between 59.4 and



Optica Publishing Group. The effect diagram of the device is shown above, and the schematic diagram is shown below. **c** The plot of transmittance vs. electric field for PDLC with different dye concentrations and the scattering effect of PDLC thin films for LC droplets with different diameters. Reproduced with permission from ref. ¹⁶⁷. Copyright 2022, Elsevier. **d** Graph of transmittance vs. time, electrode design of dye-doped PSLC, and schematic diagram of the scattering principle

41.2% by temperature, as shown in Fig. 16b. In addition, tungsten bronze (Cs_xWO_3) NRs were doped into this LC system to obtain a multi-response soft matter composite smart film; more than 95% of the NIR irradiation from 800 to 2500 nm was screened¹⁷⁶. Furthermore, Zhang¹⁷⁷

et al. prepared multiscale smart windows by mixing ATO NPs into PSLC films. An optical modulator with wavelength range from 380-nm to 5500-nm was created using PSLC visible light scattering and semiconductor nanoparticle absorption. In the visible region, the



transmittance was changed from highly transparent (78.5%) to strong light scattering (10%), while up to 80.75% of infrared invisible light was effectively shielded, as shown in Fig. 16c.

All of the above examples used electronic equipment to provide light protection. Researchers studied thermally controlled light protection equipment to save even more energy. Kragt¹⁷⁸ et al. reported a novel LC polymer-based device for light protection that reversibly increased reflectivity upon heating, as shown in Fig. 17a. Kakiu-chida¹⁷⁹ et al. developed a polymer network liquid crystal (PNLC) whose scattering intensity varied with temperature. As shown in Fig. 17b, from 20 °C to 50 °C, the direct transmittance of the PNLC decreased from 83.6% to 0.7%, and the transmittance in the infrared band changed by approximately 20%¹⁷⁹.

Researchers also developed light protection devices that responded to electricity, heat, and light¹⁸⁰. UV irradiation, temperature increase, or an electric field caused them to change them from a very transparent to a black fog state (Fig. 17c). Notably, unlike the above devices, it was metastable, meaning that an electric field was not required to maintain the black fog state; this further reduced energy consumption.

Long-term light protection strategies

Unlike optical windows used on vehicles, windows in buildings are in fixed positions and exposed to sunlight for extended periods of time. Moreover, the areas of windows in buildings are often much larger than those of optical windows in vehicles. These different usage scenarios put forward new requirements for light protection devices for buildings: (1) low-cost and large-area fabrication, (2) long-term operation with low energy consumption, and (3) capability for infrared adjustment.

The commercial P-LC materials mentioned in Section 'Scattering-based light protection' are the preferred materials. For low-cost large-area fabrication, some researchers have studied the use of flexible substrates (e.g., polyethylene terephthalate or PET) instead of rigid substrates (glass) to prepare P-LC materials. Yang¹⁸¹ et al. reported a coexistence system of polymer-dispersed and polymer-stabilized liquid crystals (PD&SLCs) by sandwiching the LC mixture into two pieces ITO-coated flexible substrates. This system can be used in large-area manufacture (with a size of $1.8 \text{ m} \times 1.1 \text{ m}$) of the smart films. In addition, the film automatically adjusted its transparency according to temperature without additional energy input. The LC droplets in the film underwent a phase transition at approximately 42 °C, the specular transmittance of visible light decreased from 80% to 1% without the need for electric field maintenance; the film was transparent at room temperature (Fig. 18a,b). Meanwhile, large-area PD&SLC films were prepared in this study; they displayed great potential for applications. In addition, PDLC films with tITO NCs on flexible substrates were investigated. Due to the localized surface plasmon resonance of ITO NCs, more than 85% of the infrared rays (750-2500 nm) were effectively shielded. The specular transmittance of visible light by the film decreased from 78% to 1.5% (Fig. 18c,d) as the result of temperature stimulation or application of an electric field²¹. Furthermore, large-area films with dimensions of $1.8 \text{ m} \times 1.1 \text{ m}$ (Fig. 18e) were prepared in this study.



in a black fog state (III). Reproduced with permission from ref. ¹⁷³. Copyright 2019, Elsevier

Huang¹⁵³ et al. applied their expertize in flexible electrodes to design and optimize a $TiO_2/Ag(Cu)/TiO_2$ (TCAT) electrode to simultaneously tune visible light transmission and near-infrared heat shielding (Fig. 19a,b).

After applying an electric field, the specular transmittance of the TCAT-based PDLC device increased from approximately 1% to 84.9%. Additionally, TCAT-based devices provided better thermal insulation than ITO-



 -doped in a PSLC smart film with on/off states a of Chemistry

based devices and good modulation performance in the 400–1600 nm band. In addition to utilizing light energy transformation and scattering transfer methods for light protection, the method of reflected transfer of light energy was used for light protection in the architectural field. Michael¹⁸² et al. presented a method for fabricating surface-tethered polymer networks (STPN) composed of helical polymer structures, where STPN acted as a structurally chiral scaffold that forced the swollen LC mixture to assume the same handedness as the polymeric material and reflect light according to the as-fabricated pitch (Fig. 19c,d). Backfilling an STPN cell with a thermally tunable SmA* mixture allowed for temperature-induced and nearly complete reflection.

A limitation of conventional light protection systems for light energy conversion is that most of the absorbed energy is eventually rereleased as localized heat, thereby causing a temperature rise, which is not good for equipment that operates for long periods of time and can cause damage indoors. A more advanced light energy conversion concept is the luminescent solar concentrator (LSC). LSCs use dyes embedded in polymer or glass plates that function as windows. These dyes absorb near-infrared light and subsequently fluoresce at longer wavelengths (Fig. 20a). A fraction of this reemitted light is trapped in a higher refractive index polymer or glass panel, which acts as a light guide. The trapped light is transported via total internal reflection



Fig. 17 Non-electrically switchable optical devices. a The relationship between transmittance and wavelength at different temperatures and the schematic diagram of device adjustment. Reproduced with permission from ref. ¹⁷⁸. Copyright 2021, Wiley Online Library. **b** Relationship between transmittance and wavelength at different temperatures and sample effect diagram. Reproduced with permission from ref. ¹⁷⁹. Copyright 2021, American Chemical Society. **c** Schematic diagram of the regulation principle of light and heat dual stimulation and the sample effect diagram. Reproduced with permission from ref. ¹⁸⁰. Copyright 2018, Wiley Online Library.



Fig. 18 Light protection strategies for buildings. a Graph of the transmittance of the sample as a function of temperature. **b** The prepared largearea films and their bending properties. **a**, **b** Reproduced with permission from ref.¹⁸¹. Copyright 2017, American Chemical Society. **c** Graph of the transmittance of the sample as a function of temperature. **d** Graph of the transmittance of the sample as a function of wavelength for different voltages. **e** The prepared large-area films and their effects at different voltages. Reproduced with permission from ref.²¹. Copyright 2017, Royal Society of Chemistry



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and only exits at the sides of the window, where it may be converted to electricity via attached photovoltaic cells^{177,181}. LSCs have not yet been widely commercialized, primarily owing to their modest efficiencies. The first loss in the LSC originates from the light emitted by the luminophore at an angle such that it is refracted out of the waveguide through an escape cone rather than reflected internally (Fig. 20b)¹⁸³. Organic luminophores are often dichroic in terms of absorption and transmission. Aligning organic dyes perpendicular to the plane of the waveguide surface (homeotropically) using a host LC leads to emissions in the plane of the waveguide, which results in a sharp decrease in the surface loss to less than 10%. Yu Xia¹⁸⁴ et al. fabricated a very transparent (79%) light protection device by coupling a multiresponse liquid crystal/polymer composite (LCPC) and planar semitransparent perovskite solar cell (ST-PSC) capable of responding (Fig. 20c-e). The device can respond to temperature or electric field stimuli and generate photoelectric conversion, and it has potential for large-area fabrication.

Conclusions and perspectives

Due to the unique thermodynamic properties of LC materials, optical technologies based on liquid crystal materials have developed rapidly over the past several decades. In this review, we provide a comprehensive evaluation of light protection applications for switchable optics based on LCs. In this context, "light protection" refers to the prevention or weakening of damage, discomfort or interference caused by inappropriate light radiation on people and light-sensitive instruments. According to the source, light radiation can be classified as natural light or artificial light. In this review, we discuss their characteristics, such as frequency, energy and irradiation area, and summarize the characteristics of light protection. From the perspective of light energy, there are three types of photoprotection principles for LCs: (1) conversion of incident light energy to other forms of energy; (2) transfer of incident light energy through reflected paths; and (3) transfer of incident light energy through scattering paths. We discuss the characteristics, advantages and applications of each of these three types of



light protection principles for light protection devices. However, in the presence of complex optical environments, comprehensive protection often requires that multiple mechanisms are coupled. We propose recommendations for light protection strategies in different light environments. When applying light protection devices to vehicles, it is necessary to focus on transient light protection strategies, while for buildings, long-term light protection strategies are appropriate.

Although several types of light protection devices for LCs were studied, those for practical applications are still very limited. Challenges and opportunities coexist in the course of accelerating the maturation of this exiting field. First, it is challenging to develop LC-based light protection devices that adapt to different external environments during longterm stable operation. Prolonged exposure to UV light makes polar groups in LC molecules susceptible to oxidation and molecular structures fracture. It also induces the fracture of organic polymers in P-LC materials, cracks in devices and photodegradation of dichroic dye molecules^{185–187}. However, LCs are highly susceptible to ion production under electrical stimulation, and these ions affect the performance of the devices in different ways and interfere with the dispersion stability of NPs in LCs, which greatly limits the lifetime of relevant devices^{188–193}. Therefore, mature light protection devices for LCs should have high UV radiation tolerance and electrochemical stability. Second, in current commercial installations, there is a lack of LC-based light protection devices that flexibly adapt to protective requirements for different external environments. Increasingly complex optical environments require a new generation of light protection devices that can operate at multiple wavelengths and react quickly, especially for passive protection. Currently, common external stimuli that cause changes in the optical performance of smart windows include electric fields, temperature, and light. Electrical stimuli correspond to active light protection devices, while thermal and optical stimuli correspond to passive light protection devices. However, the slow response to thermal stimuli and the limited wavelengths of light response limit the application of these passive light protection devices. As near-infrared light is an important component of sunlight and an important source of thermal radiation, further research can focus on developing light stimulation response devices at this range of wavelengths. Additionally, intelligent passive light protection devices should dynamically sense the light environment and adjust their optical properties according to the external light intensity. Third, these technologies face the challenge of ensuring good stability and homogeneity for extensive commercial applications. For example, optically and chemically pure materials are essential for the proper operation of Ch-LC-based devices, as any impurity can adversely affect the self-assembly of molecules¹⁹⁴. In addition, many commercially available optical windows are irregularly curved rather than planar. There are challenges in the preparation of such conformal light-proof devices. In addition, there is a broad commercial market for retrofitting conventional (traditional) windows with new windows with light protection features. Therefore, the development of deformable, flexible light protection devices will be ideal for commercial applications in the future.

There is no doubt that with the development of lighting technology, the need for light-protection devices is becoming increasingly urgent. There is strong commercial interest from related companies, such as Merck Chemicals, BASF, Dainippon Ink and other leaders in the chemical industry, where researchers are currently working on the development of light-protection devices. We expect that thorough research on a new generation of intelligent optical limiting materials will address the above issues. Programmable flexibility and self-adaptation will be critical in realizing optical materials for new demands. This system will not only provide protection from multiple wavelength bands but also respond quickly in complex environments, protect across the entire UV-visible-infrared spectrum, supply high-strength protection in some bands, and have long service life. The system should work well provided that the device is stable under high-power incident light and consumes small amounts of energy in the long-term protective state.

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Conflict of interest

The authors declare no conflicts of interest.

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References

- Cinzano, P., Falchi, F. & Elvidge, C. D. The first world atlas of the artificial night sky brightness. *Mon. Not. R. Astron. Soc.* **328**, 689–707 (2001).
- Hölker, F. et al. The dark side of light: a transdisciplinary research agenda for light pollution policy. *Ecol. Soc.* 15, 13 (2010).
- Hölker, F. et al. Light pollution as a biodiversity threat. Trends Ecol. Evol. 25, 681–682 (2010).
- Wang, L. Self-activating liquid crystal devices for smart laser protection. *Liq. Cryst.* 43, 2062–2078 (2016).
- Gosling, D. B., O'Hagan, J. B. & Quhill, F. M. Blue laser induced retinal injury in a commercial pilot at 1300 ft. Aerosp. Med. Hum. Perform. 87, 69–70 (2016).
- Liu, Z. W., Zhang, B. & Chen, Y. Recent progress in two-dimensional nanomaterials for laser protection. *Chemistry* 1, 17–43 (2019).
- Williamson, C. A. & McLin, L. N. Determination of a laser eye dazzle safety framework. J. Laser Appl. 30, 032010 (2018).
- Goto, E. et al. Impaired functional visual acuity of dry eye patients. Am. J. Ophthalmol. 133, 181–186 (2002).
- Marshall, J., O'Hagan, J. B. & Tyrer, J. Eye hazards of laser 'pointers' in perspective. Br. J. Ophthalmol. 100, 583–584 (2016).
- Murata, K. et al. Central nervous system effects and visual fatigue in VDT workers. Int. Arch. Occup. Environ. Health 63, 109–113 (1991).
- Morita, Y. et al. Effect of heat-killed *Lactobacillus paracasei* KW3110 ingestion on ocular disorders caused by visual display terminal (VDT) loads: a randomized, double-blind, placebo-controlled parallel-group study. *Nutrients* 10, 1058 (2018).
- Vila, N. et al. Blue-light filtering alters angiogenic signaling in human retinal pigmented epithelial cells culture model. *BMC Ophthalmol.* 17, 198 (2017).
- Ouyang, X. L. et al. Mechanisms of blue light-induced eye hazard and protective measures: a review. *Biomed. Pharmacother.* 130, 110577 (2020).

- 14. Xing, Q. et al. Structures of chaperone-substrate complexes docked onto the export gate in a type III secretion system. *Nat. Commun.* 9, 1773 (2018).
- Barrios, D. et al. Toward a quantitative model for suspended particle devices: optical scattering and absorption coefficients. *Sol. Energy Mater. Sol. Cells* 111, 115–122 (2013).
- Lampert, C. M. Large-area smart glass and integrated photovoltaics. Sol. Energy Mater. Sol. Cells 76, 489–499 (2003).
- 17. Cupelli, D. et al. Self-adjusting smart windows based on polymer-dispersed liquid crystals. *Sol. Energy Mater. Sol. Cells* **93**, 2008–2012 (2009).
- Khaligh, H. H. et al. Silver nanowire transparent electrodes for liquid crystalbased smart windows. *Sol. Energy Mater. Sol. Cells* **132**, 337–341 (2015).
- Khandelwal, H. et al. Electrically switchable polymer stabilised broadband infrared reflectors and their potential as smart windows for energy saving in buildings. *Sci. Rep.* 5, 11773 (2015).
- Kwon, H. K et al. Optically switchable smart windows with integrated photovoltaic devices. *Adv. Energy Mater.* 5, 1401347 (2015).
- 21. Liang, X. et al. A temperature and electric field-responsive flexible smart film with full broadband optical modulation. *Mater. Horiz.* **4**, 878–884 (2017).
- Murray, J., Ma, D. K. & Munday, J. N. Electrically controllable light trapping for self-powered switchable solar windows. ACS Photonics 4, 1–7 (2017).
- Sun, Y. J. et al. Effects of functionality of thiol monomer on electro-optical properties of polymer-dispersed liquid crystal films. *Liq. Cryst.* 44, 1086–1092 (2017).
- Wang, L., Urbas, A. M. & Li, Q. Nature-inspired emerging chiral liquid crystal nanostructures: from molecular self-assembly to DNA mesophase and nanocolloids. *Adv. Mater.* 32, 1801335 (2020).
- Wang, L. et al. Stimuli-directed self-organized chiral superstructures for adaptive windows enabled by mesogen-functionalized graphene. *Mater. Today* 20, 230–237 (2017).
- Moon, H. C. et al. Multicolored, low-power, flexible electrochromic devices based on ion gels. ACS Appl. Mater. Interfaces 8, 6252–6260 (2016).
- Oh, H. et al. Voltage-tunable multicolor, sub-1.5 V, flexible electrochromic devices based on ion gels. ACS Appl. Mater. Interfaces 9, 7658–7665 (2017).
- Runnerstrom, E. L. et al. Nanostructured electrochromic smart windows: traditional materials and NIR-selective plasmonic nanocrystals. *Chem. Commun.* 50, 10555–10572 (2014).
- Yeh, M. H. et al. Motion-driven electrochromic reactions for self-powered smart window system. ACS Nano 9, 4757–4765 (2015).
- Wang, L. & Li, Q. Photochromism into nanosystems: towards lighting up the future nanoworld. *Chem. Soc. Rev.* 47, 1044–1097 (2018).
- Wang, Y., Runnerstrom, E. L. & Milliron, D. J. Switchable materials for smart windows. Annu. Rev. Chem. Biomolecular Eng. 7, 283–304 (2016).
- Wu, L. Y. L. et al. Sol-gel based photochromic coating for solar responsive smart window. *Surf. Coat. Technol.* **320**, 601–607 (2017).
- Gao, Y. F. et al. Nanoceramic VO₂ thermochromic smart glass: a review on progress in solution processing. *Nano Energy* 1, 221–246 (2012).
- Kim, H. et al. Flexible thermochromic window based on hybridized VO₂/ graphene. ACS Nano 7, 5769–5776 (2013).
- Park, J. Y. et al. PDMS-paraffin/graphene laminated films with electrothermally switchable haze. *Carbon* 96, 805–811 (2016).
- Singh, A. K. et al. Fabrication of solar and electrically adjustable large area smart windows for indoor light and heat modulation. J. Mater. Chem. C. 5, 5917–5922 (2017).
- Zhang, J. S. et al. Hydrothermal growth of VO₂ nanoplate thermochromic films on glass with high visible transmittance. *Sci. Rep.* 6, 27898 (2016).
- Jensen, J. et al. Development and manufacture of polymer-based electrochromic devices. Adv. Funct. Mater. 25, 2073–2090 (2015).
- Yang, G. J. et al. Advances in nanomaterials for electrochromic devices. *Chem. Soc. Rev.* 49, 8687–8720 (2020).
- Kamalisarvestani, M. et al. Performance, materials and coating technologies of thermochromic thin films on smart windows. *Renew. Sustain. Energy Rev.* 26, 353–364 (2013).
- 41. Popov, N. et al. Thermotropic liquid crystal-assisted chemical and biological sensors. *Materials* **11**, 20 (2018).
- 42. Pelzl, G. & Weissflog, W. in Thermotropic Liquid Crystals: Recent Advances (ed. Ramamoorthy, A.) (Springer, 2007).
- Bisoyi, H. K. & Li, Q. Liquid crystals: versatile self-organized smart soft materials. Chem. Rev. 122, 4887–4926 (2022).
- Besharat, F., Dehghan, A. A. & Faghih, A. R. Empirical models for estimating global solar radiation: a review and case study. *Renew. Sustain. Energy Rev.* 21, 798–821 (2013).

- Omer, A. M. Energy, environment and sustainable development. *Renew. Sustain. Energy Rev.* 12, 2265–2300 (2008).
- Rathod, A. P. S., Mittal, P. & Kumar, B. Analysis of factors affecting the solar radiation received by any region. 2016 International Conference on Emerging Trends in Communication Technologies (ETCT), Dehradun, 1–4 (IEEE, 2016).
- Juzeniene, A. et al. Solar radiation and human health. *Rep. Prog. Phys.* 74, 066701 (2011).
- Sengupta, M. et al. The national solar radiation data base (NSRDB). *Renew. Sustain. Energy Rev.* 89, 51–60 (2018).
- Behar-Cohen, F. et al. Light-emitting diodes (LED) for domestic lighting: any risks for the eye? *Prog. Retinal Eye Res.* 30, 239–257 (2011).
- Tao, J. X., Zhou, W. C. & Zhu, X. G. Mitochondria as potential targets and initiators of the blue light hazard to the retina. *Oxid. Med. Cell. Longev.* 2019, 6435364 (2019).
- Sen, Z. Solar Energy Fundamentals and Modeling Techniques: Atmosphere, Environment, Climate Change and Renewable Energy (Springer Science & Business Media, 2008).
- Dinh, T. V. et al. A review on non-dispersive infrared gas sensors: Improvement of sensor detection limit and interference correction. *Sens. Actuators B: Chem.* 231, 529–538 (2016).
- Ogawa, T. et al. Pedestrian detection and tracking using in-vehicle lidar for automotive application. 2011 IEEE Intelligent Vehicles Symposium (IV). Baden-Baden, Germany, 734–739 (IEEE, 2011).
- Rogalski, A., Antoszewski, J. & Faraone, L. Third-generation infrared photodetector arrays. J. Appl. Phys. 105, 091101 (2009).
- Shao, X. M. et al. The influence of sunlight irradiation on the characteristics of InGaAs detectors. Proceedings of SPIE 9220, Infrared Sensors, Devices, and Applications IV. San Diego, California, 83–88 (SPIE, 2014).
- Silberman, N. & Fergus, R. Indoor scene segmentation using a structured light sensor. 2011 IEEE International Conference on Computer Vision Workshops (ICCV Workshops). Barcelona, Spain, 601–608 (IEEE, 2011).
- 57. Loonen, R. C. G. M. et al. Climate adaptive building shells: State-of-the-art and future challenges. *Renew. Sustain. Energy Rev.* **25**, 483–493 (2013).
- Abbondanzieri, E. A. et al. Direct observation of base-pair stepping by RNA polymerase. *Nature* 438, 460–465 (2005).
- Smith, S. B., Cui, Y. J. & Bustamante, C. Overstretching B-DNA: the elastic response of individual double-stranded and single-stranded DNA molecules. *Science* 271, 795–799 (1996).
- Webb, C. & Jones, J. Handbook of Laser Technology and Applications (Three-Volume Set) (CRC Press, 2003).
- Pert, G. J. Output characteristics of amplified-stimulated-emission lasers. J. Optical Soc. Am. B 11, 1425–1435 (1994).
- 62. Federal Aviation Administration at https://www.faa.gov/about/initiatives/ lasers/laws (2021).
- 63. TNO. The Outdoor Use of Lasers and Other High Intensity Light Sources in Relation to Air Traffic Safety (TNO, 2009).
- Thomas, R. J. et al. A procedure for laser hazard classification under the Z136.1-2000 American National Standard for Safe Use of Lasers. J. Laser Appl. 14, 57–66 (2002).
- Smith, J. F. et al. Are you a laser product manufacturer?: Interpretation of the FDA regulations applicable to laser product manufacturers. J. Laser Appl. 13, 150–153 (2001).
- Schulmeister, K. The new edition of the international laser product safety standard IEC 60825-1 (White paper) (2017).
- Svensson, S. et al. Countering laser pointer threats to road safety. Proceedings of SPIE 6402, Optics and Photonics for Counterterrorism and Crime Fighting II. Stockholm, Sweden, 41–48 (SPIE, 2006).
- Yang D. K, & Wu S. T. Fundamentals of Liquid Crystal Devices (John Wiley & Sons, 2014).
- Jiang, Y. F., Shin, Y. & Yang, D. K. Dual-mode switchable liquid-crystal window. *Phys. Rev. Appl.* 12, 054037 (2019).
- Dierking, I. Science of the present meets the life of the past. *Liq. Cryst. Today* 25, 10–11 (2016).
- Foglia, N. O. et al. Role of core electrons in quantum dynamics using TDDFT. J. Chem. Theory Comput. 13, 77–85 (2017).
- Seki, H., Uchida, T. & Shibata, Y. Dichroic dyes for guest-host liquid-crystal cells. *Mol. Cryst. Liq. Cryst.* 138, 349–365 (1986).
- Heilmeier, G. H. & Zanoni, L. A. Guest-host interactions in nematic liquid crystals. *Appl. Phys. Lett.* 13, 91–92 (1968).
- Dorozhkina, G. N. et al. New PDLC-films doped by black dye. Mol. Cryst. Liq. Cryst. Sci. Technol. Sect. A. Mol. Cryst. Liq. Cryst. 367, 277–286 (2001).

- Heppke, G. et al. Colored and black liquid crystalline mixtures with new antraquinone dyes. *Mol. Cryst. Liq. Cryst.* 94, 191–204 (1983).
- Olifierczuk, M. & Zielinski, J. The contrast ratio in twisted nematic with black dichroic dye LCD. Synth. Met. 109, 223–227 (2000).
- Prudnikova, E., Umanskii, B. & Plyusnina, T. Synthesis of new dichroic dyes with negative dichroism for a black mixture. *Mol. Cryst. Liq. Cryst. Sci. Technol. Sect. A. Mol. Cryst. Liq. Cryst.* 332, 37–41 (1999).
- 78. Bahadur, B. Liquid Crystals: Applications and Uses (World Scientific, 1990).
- Deshmukh, R. R. & Malik, M. K. Effect of dichroic dye on phase separation kinetics and electro-optical characteristics of polymer dispersed liquid crystals. J. Phys. Chem. Solids 74, 215–224 (2013).
- Hsiao, Y. C. et al. Preparation and characterization of PDLC films formed using a two-step procedure. *Adv. Polym. Technol.* 26, 14–20 (2007).
- Kłosowicz, S. J. & Aleksander, M. Effect of polymer-dispersed liquid crystal morphology on its optical performance. *Opto-Electron. Rev.* **12**, 305–312 (2004).
- Kumar, P., Sharma, V. & Raina, K. K. Studies on inter-dependency of electrooptic characteristics of orange azo and blue anthraquinone dichroic dye doped polymer dispersed liquid crystals. *J. Mol. Liq.* 251, 407–416 (2018).
- Li, W. B. et al. Effects of the structures of polymerizable monomers on the electro-optical properties of UV cured polymer dispersed liquid crystal films. J. Polym. Sci. B Polym. Phys. 46, 1369–1375 (2008).
- Malik, P. & Raina, K. K. Dichroic dye-dependent studies in guest-host polymer-dispersed liquid crystal films. *Phys. B Condens. Matter* **405**, 161–166 (2010).
- West, J. L., Ondris-Crawford, R. & Erdmann, M. A. Dichroic dye containing polymer-dispersed liquid crystal films. Proceedings of SPIE 1257, Liquid Crystal Displays and Applications. Santa Clara, CA, 76–83 (SPIE, 1990).
- Huh, J. W. et al. lon-doped liquid-crystal cell with low opaque-state specular transmittance based on electro-hydrodynamic effect. *Dyes Pigments* 150, 16–20 (2018).
- Li, B. X. et al. Dye-doped dual-frequency nematic cells as fast-switching polarization-independent shutters. *Opt. Express* 27, 3861–3866 (2019).
- Yu, B. H. et al. Light shutter using dichroic-dye-doped long-pitch cholesteric liquid crystals. Opt. Express 21, 29332–29337 (2013).
- Hartland, G. V. et al. What's so hot about electrons in metal nanoparticles? ACS Energy Lett. 2, 1641–1653 (2017).
- Tesler, A. B. et al. Tunable localized plasmon transducers prepared by thermal dewetting of percolated evaporated gold films. J. Phys. Chem. C. 115, 24642–24652 (2011).
- Yablonovitch, E. & Cody, G. D. Intensity enhancement in textured optical sheets for solar cells. *IEEE Trans. Electron Devices* 29, 300–305 (1982).
- Brochard, F. & De Gennes, P. G. Theory of magnetic suspensions in liquid crystals. J. Phys. 31, 691–708 (1970).
- Bukowczan, A., Hebda, E. & Pielichowski, K. The influence of nanoparticles on phase formation and stability of liquid crystals and liquid crystalline polymers. *J. Mol. Liq.* **321**, 114849 (2021).
- Gao, R. Q. L. et al. Antimony doped tin oxide infrared shielding films for cooling silicon solar cells. in Advanced Functional Materials (ed. Han, Y. F.) 817–829 (Springer, 2018).
- Garcia, G. et al. Dynamically modulating the surface Plasmon resonance of doped semiconductor nanocrystals. *Nano Lett.* 11, 4415–4420 (2011).
- Tan, Q. et al. Preparation of stable aqueous suspensions of antimony-doped tin oxide nanoparticles used for transparent and thermal insulation fluorocarbon coating. *Colloid Polym. Sci.* **292**, 3233–3241 (2014).
- 97. Kim, H. N. & Yang, S. Responsive smart windows from nanoparticle–polymer composites. *Adv. Funct. Mater.* **30**, 1902597 (2020).
- Jiang, G. Q. et al. Gold nanorods dispersed in homopolymer films: optical properties controlled by self-assembly and percolation of nanorods. ACS Nano 6, 1578–1588 (2012).
- Liu, Q. K. et al. Self-alignment of plasmonic gold nanorods in reconfigurable anisotropic fluids for tunable bulk metamaterial applications. *Nano Lett.* 10, 1347–1353 (2010).
- Stebe, K. J., Lewandowski, E. & Ghosh, M. Oriented assembly of metamaterials. Science 325, 159–160 (2009).
- Xue, C. M. et al. Photomodulated self-assembly of hydrophobic thiol monolayer-protected gold nanorods and their alignment in thermotropic liquid crystal. J. Phys. Chem. C 117, 21603–21608 (2013).
- Moran, M. J. et al. Driving a liquid crystal phase transition using a photochromic hydrazone. J. Am. Chem. Soc. 140, 13623–13627 (2018).

- Sheetah, G. H. et al. Electric switching of visible and infrared transmission using liquid crystals co-doped with plasmonic gold nanorods and dichroic dyes. *Opt. Express* 26, 22264–22272 (2018).
- Manias, E. et al. Polypropylene/Montmorillonite Nanocomposites. Review of the synthetic routes and materials properties. *Chem. Mater.* 13, 3516–3523 (2001).
- Stelzig, S. H., Klapper, M. & Müllen, K. A simple and efficient route to transparent nanocomposites. *Adv. Mater.* 20, 929–932 (2008).
- Wen, J. Y. & Wilkes, G. L. Organic/inorganic hybrid network materials by the sol-gel approach. *Chem. Mater.* 8, 1667–1681 (1996).
- Choudhary, A., Singh, G. & Biradar, A. M. Advances in gold nanoparticle–liquid crystal composites. *Nanoscale* 6, 7743–7756 (2014).
- 108. Prakash, J. et al. Metal oxide-nanoparticles and liquid crystal composites: a
- review of recent progress. J. Mol. Liq. **297**, 112052 (2020). 109. Friedel, G. Les états mésomorphes de la matière. Ann. Phys. **9**, 273–474 (1922).
- Hieuer, G. Les etals mesoniorpries de la materie. Ann. Phys. 9, 273–474 (1922)
 Kelker, H. History of liquid crystals. Mol. Cryst. Liq. Cryst. 21, 1–48 (1973).
- Broer, D. J. et al. Photo-induced diffusion in polymerizing chiral-nematic media. Adv. Mater. 11, 573–578 (1999).
- Mulder, D. J., Schenning, A. P. H. J. & Bastiaansen, C. W. M. Chiral-nematic liquid crystals as one dimensional photonic materials in optical sensors. *J. Mater. Chem. C.* 2, 6695–6705 (2014).
- Ha, N. Y. et al. Fabrication of a simultaneous red–green–blue reflector using single-pitched cholesteric liquid crystals. *Nat. Mater.* 7, 43–47 (2008).
- Huang, Y. H., Zhou, Y. & Wu, S. T. Broadband circular polarizer using stacked chiral polymer films. *Opt. Express* 15, 6414–6419 (2007).
- Binet, C., Mitov, M. & Boudet, A. Bragg reflections in cholesteric liquid crystals: from selectivity to broadening and reciprocally. *Mol. Cryst. Liq. Cryst. Sci. Technol. Sect. A. Mol. Cryst. Liq. Cryst.* **339**, 111–123 (2000).
- Boudet, A. et al. Microstructure of variable pitch cholesteric films and its relationship with the optical properties. *Eur. Phys. J. E* 2, 247–253 (2000).
- Mitov, M., Boudet, A. & Sopéna, P. From selective to wide-band light reflection: a simple thermal diffusion in a glassy cholesteric liquid crystal. *Eur. Phys. J. B-Condens. Matter Complex Syst.* 8, 327–330 (1999).
- Zografopoulos, D. C. et al. Theoretical and experimental optical studies of cholesteric liquid crystal films with thermally induced pitch gradients. *Phys. Rev. E* 73, 061701 (2006).
- Li, Y. N. Q. et al. Broadband cholesteric liquid crystal lens for chromatic aberration correction in catadioptric virtual reality optics. *Opt. Express* 29, 6011–6020 (2021).
- Hikmet, R. A. M. & Kemperman, H. Electrically switchable mirrors and optical components made from liquid-crystal gels. *Nature* 392, 476–479 (1998).
- Broer, D. J., Lub, J. & Mol, G. N. Photo-controlled diffusion in reacting liquid crystals: a new tool for the creation of complex molecular architectures. *Macromol. Symposia* **117**, 33–42 (1997).
- Broer, D. J., Lub, J. & Mol, G. N. Wide-band reflective polarizers from cholesteric polymer networks with a pitch gradient. *Nature* 378, 467–469 (1995).
- Broer, D. J. Deformed chiral-nematic networks obtained by polarized excitation of a dichroic photoinitiator. *Curr. Opin. Solid State Mater. Sci.* 6, 553–561 (2002).
- Lub, J. & Broer, D. J. & Van de Witte, P. Colourful photo-curable coatings for application in the electro-optical industry. *Prog. Org. Coat.* 45, 211–217 (2002).
- Lub, J. et al. Formation of optical films by photo-polymerisation of liquid crystalline acrylates and application of these films in liquid crystal display technology. *Mol. Cryst. Liq. Cryst.* **429**, 77–99 (2005).
- Hu, X. W. et al. Pitch gradation by ion-dragging effect in polymer-stabilized cholesteric liquid crystal reflector device. *Polymers* 12, 96 (2020).
- 127. Guo, J. B. et al. Effect of network concentration on the performance of polymer-stabilized cholesteric liquid crystals with a double-handed circularly polarized light reflection band. J. Phys. Chem. C. 113, 16538–16543 (2009).
- Jain, A. K. & Deshmukh, R. R. An overview of polymer-dispersed liquid crystals composite films and their applications. in Liquid Crystals and Display Technology (eds Ghamsari, M. S. & Carlescu, I.) 1–68 (IntechOpen, 2020).
- Doane, J. W. et al. Field controlled light scattering from nematic microdroplets. *Appl. Phys. Lett.* 48, 269–271 (1986).
- Saeed, M. H. et al. Recent advances in the polymer dispersed liquid crystal composite and its applications. *Molecules* 25, 5510 (2020).
- Li, W. B. et al. Control of the microstructure of polymer network and effects of the microstructures on light scattering properties of UV-cured polymer-dispersed liquid crystal films. J. Polym. Sci. B Polym. Phys. 46, 2090–2099 (2008).
- 132. Drzaic, P. S. Reorientation dynamics of polymer dispersed nematic liquid crystal films. *Liq. Cryst.* **3**, 1543–1559 (1988).

- 133. Nicoletta, F. P. et al. Changes in the electro-optical properties of a PDLC system by monomer substitution. *Mol. Cryst. Liq. Cryst. Sci. Technol. Sect. A. Mol. Cryst. Liq. Cryst.* **336**, 93–99 (1999).
- Wu, B. G., Erdmann, J. H. & Doane, J. W. Response times and voltages for PDLC light shutters. *Liq. Cryst.* 5, 1453–1465 (1989).
- Zhang, Y. et al. Effects of crosslinking agent/diluents/thiol on morphology of the polymer matrix and electro-optical properties of polymer-dispersed liquid crystal. *Liq. Cryst.* **45**, 728–735 (2018).
- Li, C. X. et al. A study on the polymer structures and electro-optical properties of epoxy-mercaptan-based polymer dispersed liquid crystal films. *Liq. Cryst.* 46, 1718–1726 (2019).
- Ahmad, F., Luqman, M. & Jamil, M. Advances in the metal nanoparticles (MNPs) doped liquid crystals and polymer dispersed liquid crystal (PDLC) composites and their applications-a review. *Mol. Cryst. Liq. Cryst.* **731**, 1–33 (2021).
- Ji, Y. Y. et al. Terahertz birefringence anisotropy and relaxation effects in polymer-dispersed liquid crystal doped with gold nanoparticles. *Opt. Express* 28, 17253–17265 (2020).
- Mishra, K. K., Dubey, S. K. & Mani, S. A. Optical characterization of inorganic nanoparticles doped in polymer dispersed liquid crystal. *Mol. Cryst. Liq. Cryst.* 647, 244–252 (2017).
- Wu, Q. Q. & Wang, Y. Q. Low driving voltage ITO doped polymer-dispersed liquid crystal film and reverse voltage pulse driving method. *Appl. Opt.* 56, 8159–8163 (2017).
- Ma, Y. D., Wu, B. G. & Xu, G. Reverse-mode microdroplet liquid crystal display. Proceedings of SPIE 1257, Liquid Crystal Displays and Applications. Santa Clara, CA, 46–57 (SPIE, 1990).
- Meng, C. L. et al. Normally transparent smart window with haze enhancement via inhomogeneous alignment surface. *Lig. Cryst.* 46, 484–491 (2019).
- Deng, Y. et al. High-efficiency responsive smart windows fabricated by carbon nanotubes modified by liquid crystalline polymers. *Crystals* **11**, 440 (2021).
- Kakiuchida, H., Matsuyama, A. & Ogiwara, A. Normal-and reverse-mode thermoresponsive controllability in optical attenuation of polymer network liquid crystals. ACS Appl. Mater. Interfaces 11, 19404–19412 (2019).
- Heilmeier, G. H., Zanoni, L. A. & Barton, L. A. Dynamic scattering in nematic liquid crystals. *Appl. Phys. Lett.* **13**, 46–47 (1968).
- 146. Zhang, B. R. & Kitzerow, H. Pattern formation in a nematic liquid crystal mixture with negative anisotropy of the electric conductivity- a long-known system with "inverse" light scattering revisited. J. Phys. Chem. B 120, 6865–6871 (2016).
- 147. Koide, N. The Liquid Crystal Display Story: 50 Years of Liquid Crystal R&D that lead The Way to the Future (Springer Verlag, 2016).
- Bodenschatz, E., Zimmermann, W. & Kramer, L. On electrically driven patternforming instabilities in planar nematics. J. Phys. 49, 1875–1899 (1988).
- Zhan, Y. Y. et al. Electrohydrodynamic instabilities for smart window applications. Lig. Cryst. 47, 977–983 (2020).
- Zhan, Y. Y. et al. Light-driven electrohydrodynamic instabilities in liquid crystals. Adv. Funct. Mater. 28, 1707436 (2018).
- Perlmutter, S. H., Doroski, D. & Moddel, G. Degradation of liquid crystal device performance due to selective adsorption of ions. *Appl. Phys. Lett.* 69, 1182–1184 (1996).
- 152. Xu, D. M. et al. Image sticking in liquid crystal displays with lateral electric fields. *J. Appl. Phys.* **116**, 193102 (2014).
- Yoon, T. H. et al. Control of the haze value by the electro-hydrodynamic effect in a liquid crystal cell. Proceedings of SPIE 10735, Liquid Crystals XXII, 107350S (SPIE, 2018).
- 154. Špirk, S. et al. Utilization of the validated windshield material model in simulation of tram to pedestrian collision. *Materials* **14**, 265 (2021).
- Hu, X. W. et al. Stable and scalable smart window based on polymer stabilized liquid crystals. J. Appl. Polym. Sci. 137, 48917 (2020).
- Mitov, M. & Dessaud, N. Going beyond the reflectance limit of cholesteric liquid crystals. *Nat. Mater.* 5, 361–364 (2006).
- Shi, Z. Q. et al. Fabrication of dye-doped polymer-dispersed liquid crystals with low driving voltage based on nucleophile-initiated thiol-ene click reaction. *Liq. Cryst.* 45, 579–585 (2018).
- Bronnikov, S., Kostromin, S. & Zuev, V. Polymer-dispersed liquid crystals: progress in preparation, investigation, and application. J. Macromol. Sci. B 52, 1718–1735 (2013).
- Hsu, C. C. et al. Low switching voltage ZnO quantum dots doped polymerdispersed liquid crystal film. Opt. Express 24, 7063–7068 (2016).

- Hu, J. M. et al. Combined effect of hydroxylated and fluorinated acrylate monomers on improving the electro-optical and mechanical performances of PDLC-films. *Liq. Cryst.* **49**, 769–779 (2022).
- Jayoti, D., Malik, P. & Singh, A. Analysis of morphological behaviour and electro-optical properties of silica nanoparticles doped polymer dispersed liquid crystal composites. J. Mol. Liq. 225, 456–461 (2017).
- Satapathy, P. et al. Switchable smart windows using a biopolymer network of cellulose nanocrystals imposed on a nematic liquid crystal. *Appl. Phys. Lett.* 117, 103702 (2020).
- Madhuri, P. L. et al. Cochleate-doped liquid crystal as switchable metamaterial window mediated by molecular orientation modified aggregation. *Part. Part. Syst. Charact.* 37, 2000067 (2020).
- Abdulhalim, I. et al. Novel easy to fabricate liquid crystal composite with potential for electrically or thermally controlled transparency windows. *Opt. Express* 27, 17387–17401 (2019).
- Pappu, L. M. et al. Voltage controlled scattering from porous silicon Mieparticles in liquid crystals. J. Mol. Liq. 281, 108–116 (2019).
- Chen, C. W. et al. Normally transparent smart window based on electrically induced instability in dielectrically negative cholesteric liquid crystal. *Optical Mater. Express* 8, 691–697 (2018).
- Katariya-Jain, A. & Deshmukh, R. R. Effects of dye doping on electro-optical, thermo-electro-optical and dielectric properties of polymer dispersed liquid crystal films. J. Phys. Chem. Solids 160, 110363 (2022).
- Heo, J., Huh, J. W. & Yoon, T. H. Fast-switching initially-transparent liquid crystal light shutter with crossed patterned electrodes. *AIP Adv.* 5, 047118 (2015).
- Li, Q. et al. Reversible photoswitchable axially chiral dopants with high helical twisting power. J. Am. Chem. Soc. 129, 12908–12909 (2007).
- Ma, J. et al. Light-driven nanoscale chiral molecular switch: reversible dynamic full range color phototuning. *Chem. Commun.* 46, 3463–3465 (2010).
- 171. Wang, H. et al. Photochemically and thermally driven full-color reflection in a self-organized helical superstructure enabled by a halogen-bonded chiral molecular switch. *Angew. Chem. Int. Ed.* **57**, 1627–1631 (2018).
- Kim, M. et al. Fabrication of microcapsules for dye-doped polymer-dispersed liquid crystal-based smart windows. ACS Appl. Mater. Interfaces 7, 17904–17909 (2015).
- Baliyan, V. K., Jeong, K. U. & Kang, S. W. Dichroic-dye-doped short pitch cholesteric liquid crystals for the application of electrically switchable smart windows. *Dyes Pigments* **166**, 403–409 (2019).
- Madhuri, P. L. et al. Hybrid vanadium dioxide-liquid crystal tunable nonreciprocal scattering metamaterial smart window for visible and infrared radiation control. *Optical Mater. Express* **11**, 3023–3037 (2021).
- Liang, X. et al. Dual-band modulation of visible and near-infrared light transmittance in an all-solution-processed hybrid micro-nano composite film. ACS Appl. Mater. Interfaces 9, 40810–40819 (2017).
- 176. Liang, X. et al. A roll-to-roll process for multi-responsive soft-matter composite films containing Cs_xWO_3 nanorods for energy-efficient smart window applications. *Nanoscale Horiz.* **2**, 319–325 (2017).
- 177. Zhang, Z. B. et al. Visible and infrared optical modulation of PSLC smart films doped with ATO nanoparticles. *Dalton Trans.* **50**, 10033–10040 (2021).
- Kragt, A. J. J. et al. 'Smart' light-reflective windows based on temperature responsive twisted nematic liquid crystal polymers. J. Polym. Sci. 59, 1278–1284 (2021).
- Kakiuchida, H. et al. Thermoresponsive reflective scattering of meso-scale phase separation structures of uniaxially orientation-ordered liquid crystals and reactive mesogens. ACS Appl. Mater. Interfaces 13, 41066–41074 (2021).
- Oh, S. W. et al. Optical and thermal switching of liquid crystals for self-shading windows. Adv. Sustain. Syst. 2, 1700164 (2018).
- Guo, S. M. et al. Preparation of a thermally light-transmittance-controllable film from a coexistent system of polymer-dispersed and polymer-stabilized liquid crystals. ACS Appl. Mater. Interfaces 9, 2942–2947 (2017).
- McConney, M. E. et al. Thermally induced, multicolored hyper-reflective cholesteric liquid crystals. *Adv. Mater.* 23, 1453–1457 (2011).
- Debije, M. G. & Verbunt, P. P. C. Thirty years of luminescent solar concentrator research: Solar energy for the built environment. *Adv. Energy Mater.* 2, 12–35 (2012).
- Xia, Y. et al. High-efficiency and reliable smart photovoltaic windows enabled by multiresponsive liquid crystal composite films and semi-transparent perovskite solar cells. *Adv. Energy Mater.* 9, 1900720 (2019).

- Lin, P. T. et al. UV stability of high birefirngence liquid crystals. *Mol. Cryst. Liq. Cryst.* 411, 243–253 (2004).
- Wen, C. H., Gauza, S. & Wu, S. T. Ultraviolet stability of liquid crystals containing cyano and isothiocyanato terminal groups. *Liq. Cryst.* **31**, 1479–1485 (2004).
- Zhou, Y. F. et al. Harnessing the properties of colloidal quantum dots in luminescent solar concentrators. *Chem. Soc. Rev.* 47, 5866–5890 (2018).
- Garbovskiy, Y. & Glushchenko, I. Nano-objects and ions in liquid crystals: ion trapping effect and related phenomena. *Crystals* 5, 501–533 (2015).
- Garbovskiy, Y. Switching between purification and contamination regimes governed by the ionic purity of nanoparticles dispersed in liquid crystals. *Appl. Phys. Lett.* **108**, 121104 (2016).
- Garbovskiy, Y. Electrical properties of liquid crystal nano-colloids analysed from perspectives of the ionic purity of nano-dopants. *Liq. Cryst.* 43, 648–653 (2016).
- Garbovskiy, Y. Kinetics of ion-capturing/ion-releasing processes in liquid crystal devices utilizing contaminated nanoparticles and alignment films. *Nanomaterials* 8, 59 (2018).
- Narayanan, T. S. N. S. et al. Electroless nanocomposite coatings: synthesis, characteristics, and applications. in Handbook of Nanoelectrochemistry: Electrochemical Synthesis Methods, Properties, and Characterization Techniques (eds Aliofkhazraei, M. & Makhlouf, A. S. H.) 1–23 (Springer, 2016).
- Sudagar, J., Lian, J. S. & Sha, W. Electroless nickel, alloy, composite and Nano coatings-A critical review. J. Alloy. Compd. 571, 183–204 (2013).
- Poryvai, A. et al. Determination of optical purity of lactic acid-based chiral liquid crystals and corresponding building blocks by chiral high-performance liquid chromatography and supercritical fluid chromatography. *Molecules* 24, 1099 (2019).
- 195. Guo, J. B. et al. Electrothermal switching characteristics from a hydrogenbonded polymer network structure in cholesteric liquid crystals with a double-handed circularly polarized light reflection band. J. Phys. Chem. B 115, 861–868 (2011).
- Chen, F. J. et al. Novel photo-polymerizable chiral hydrogen-bonded selfassembled complexes: preparation, characterization and the utilization as a thermal switching reflective color film. J. Mater. Chem. 21, 8574–8582 (2011).
- Li, Y. N. et al. Photodynamic chiral molecular switches with thermal stability: from reflection wavelength tuning to handedness inversion of self-organized helical superstructures. *Angew. Chem. Int. Ed.* **52**, 13703–13707 (2013).
- Li, Y. N. et al. Azoarenes with opposite chiral configurations: light-driven reversible handedness inversion in self-organized helical superstructures. *Angew. Chem. Int. Ed.* 52, 8925–8929 (2013).
- Wang, L. et al. Reversible near-infrared light directed reflection in a selforganized helical superstructure loaded with upconversion nanoparticles. J. Am. Chem. Soc. 136, 4480–4483 (2014).
- Khandelwal, H. et al. Electrically tunable infrared reflector with adjustable bandwidth broadening up to 1100 nm. J. Mater. Chem. A 4, 6064–6069 (2016).
- van Heeswijk, E. P. A. et al. Humidity-gated, temperature-responsive photonic infrared reflective broadband coatings. J. Mater. Chem. A 7, 6113–6119 (2019).
- Shibaev, P. V., Schaumburg, K. & Plaksin, V. Responsive chiral hydrogenbonded polymer composites. *Chem. Mater.* 14, 959–961 (2002).
- Shibaev, P. V. et al. Color changing cholesteric polymer films sensitive to amino acids. *Macromolecules* 39, 3986–3992 (2006).
- Stumpel, J. E. et al. Stimuli-responsive materials based on interpenetrating polymer liquid crystal hydrogels. *Adv. Funct. Mater.* 25, 3314–3320 (2015).
- Hu, W. et al. Thermally controllable reflective characteristics from rupture and self-assembly of hydrogen bonds in cholesteric liquid crystals. J. Phys. Chem. B 113, 13882–13885 (2009).
- He, Z. M. et al. Silicon nanostructure-doped polymer/nematic liquid crystal composites for low voltage-driven smart windows. *Nanotechnology* 33, 085205 (2021).
- 207. Oh, S. W. et al. Optical and electrical switching of cholesteric liquid crystals containing azo dye. *RSC Adv.* **7**, 19497–19501 (2017).

- Ghosh, A. & Norton, B. Optimization of PV powered SPD switchable glazing to minimise probability of loss of power supply. *Renew. Energy* 131, 993–1001 (2019).
- Wu, W. et al. Electrochromic metal oxides: recent progress and prospect. Adv. Electron. Mater. 4, 1800185 (2018).
- Cui, Y. Y. et al. Thermochromic VO₂ for energy-efficient smart windows. *Joule* 2, 1707–1746 (2018).
- Ghosh, A., Norton, B. & Duffy, A. Measured overall heat transfer coefficient of a suspended particle device switchable glazing. *Appl. Energy* 159, 362–369 (2015).
- Ghosh, A., Norton, B. & Duffy, A. First outdoor characterisation of a PV powered suspended particle device switchable glazing. *Sol. Energy Mater. Sol. Cells* 157, 1–9 (2016).
- Ghosh, A. & Norton, B. Durability of switching behaviour after outdoor exposure for a suspended particle device switchable glazing. *Sol. Energy Mater. Sol. Cells* 163, 178–184 (2017).
- Nundy, S. & Ghosh, A. Thermal and visual comfort analysis of adaptive vacuum integrated switchable suspended particle device window for temperate climate. *Renew. Energy* **156**, 1361–1372 (2020).
- Zhao, Q. et al. Printing of WO₃/ITO nanocomposite electrochromic smart windows. Sol. Energy Mater. Sol. Cells **194**, 95–102 (2019).
- Chang, T. C. et al. Optical design and stability study for ultrahigh-performance and long-lived vanadium dioxide-based thermochromic coatings. *Nano Energy* 44, 256–264 (2018).
- Panagopoulou, M. et al. Thermochromic performance of Mg-doped VO₂ thin films on functional substrates for glazing applications. *Sol. Energy Mater. Sol. Cells* **157**, 1004–1010 (2016).
- Qian, X. K. et al. Bioinspired multifunctional vanadium dioxide: improved thermochromism and hydrophobicity. *Langmuir* **30**, 10766–10771 (2014).
- Zhan, Y. J. et al. Enhanced thermal stability and thermochromic properties of VO_x-based thin films by room-temperature magnetron sputtering. *Sol. Energy Mater. Sol. Cells* **174**, 102–111 (2018).
- Zhang, J. et al. Mesoporous SiO₂/VO₂ double-layer thermochromic coating with improved visible transmittance for smart window. *Sol. Energy Mater. Sol. Cells* 162, 134–141 (2017).
- Zhang, W. X. et al. Easily processable temperature-responsive infraredreflective polymer coatings. ACS Omega 2, 3475–3482 (2017).
- Zhou, Y. et al. Temperature-responsive hydrogel with ultra-large solar modulation and high luminous transmission for "smart window" applications. J. Mater. Chem. A 2, 13550–13555 (2014).
- Zhu, J. T. et al. Composite film of vanadium dioxide nanoparticles and ionic liquid–nickel–chlorine complexes with excellent visible thermochromic performance. ACS Appl. Mater. Interfaces 8, 29742–29748 (2016).
- Chen, Y. X. et al. VO₂/Nickel-bromine-ionic liquid composite film for thermochromic application. Sol. Energy Mater. Sol. Cells 196, 124–130 (2019).
- 225. Gao, Y. F. et al. Enhanced chemical stability of VO₂ nanoparticles by the formation of SiO₂/VO₂ core/shell structures and the application to transparent and flexible VO₂-based composite foils with excellent thermochromic properties for solar heat control. *Energy Environ. Sci.* **5**, 6104–6110 (2012).
- Lin, J. et al. Thermochromic halide perovskite solar cells. Nat. Mater. 17, 261–267 (2018).
- Lu, Q. et al. Periodic micro-patterned VO₂ thermochromic films by mesh printing. J. Mater. Chem. C. 4, 8385–8391 (2016).
- Khandelwal, H., Schenning, A. P. H. J. & Debije, M. G. Infrared regulating smart window based on organic materials. *Adv. Energy Mater.* 7, 1602209 (2017).
- Wu, S. T. Absorption measurements of liquid crystals in the ultraviolet, visible, and infrared. J. Appl. Phys. 84, 4462–4465 (1998).
- Huang, J. H. et al. Simultaneous achievement of high visible transmission and near-infrared heat shielding in flexible liquid crystal-based smart windows via electrode design. *Sol. Energy* 188, 857–864 (2019).