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Multi-stimuli-responsive luminescence enabled by crown ether anchored chiral antimony halide phosphors†

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Stimuli-responsive optical materials have provided a powerful impetus for the development of intelligent optoelectronic devices. The family of organic–inorganic hybrid metal halides, distinguished by their structural diversity, presents a prospective platform for the advancement of stimuli-responsive optical materials. Here, we have employed a crown ether to anchor the A-site cation of a chiral antimony halide, enabling convenient control and modulation of its photophysical properties. The chirality-dependent asymmetric lattice distortion of inorganic skeletons assisted by a crown ether promotes the formation of self-trapped excitons (STEs), leading to a high photoluminescence quantum yield of over 85%, concomitant with the effective circularly polarized luminescence. The antimony halide enantiomers showcase highly sensitive stimuli-responsive luminescent behaviours towards excitation wavelength and temperature simultaneously, exhibiting a versatile reversible colour switching capability from blue to white and further to orange. *In situ* temperature-dependent luminescence spectra, time-resolved luminescence spectra and theoretical calculations reveal that the multi-stimuli-responsive luminescent behaviours stem from distinct STEs within zero-dimensional lattices. By virtue of the inherent flexibility and adaptability, these chiral antimony chlorides have promising prospects for future applications in cutting-edge fields such as multifunctional illumination technologies and intelligent sensing devices.

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Introduction

Stimuli-responsive materials have gained considerable attention for their talent to toggle or alter their intrinsic properties upon external stimuli, embracing light,^{1,2} temperature,³ electric field,^{4,5} chemical perturbation,⁶ *etc.* Numerous artificial sensing devices, therefore, have been established by incorporating such stimuli-responsive materials. In particular, stimuli-responsive luminescence materials,⁷ being able to afford feedback on ambient stimuli through optical outputs in directable manners,⁸ have emerged as trending on account of their various advantages, encompassing facile detection of luminescent signals, exceptional sensitivity, rapid response speed, and an expansive scope for emission intensity and wavelength modulations.⁹ Hence, various stimuli-responsive luminescence

materials including organic small molecules,^{10,11} polymers,¹² organometallic complexes,¹³ and hybrid organic–inorganic metal halides (HOMHs)¹⁴ have been extensively utilized in prominent domains^{15–17} such as intelligent optoelectronics,^{18,19} biosensors,²⁰ and information storage.^{21–23}

Among these stimuli-responsive luminescence materials, HOMHs feature rich chemical composition and structural diversity²⁴ along with tailorable bandgaps, long carrier diffusion lengths,²⁵ large oscillator strengths and exceptional optical absorption coefficients.²⁶ Furthermore, compared to pure organic molecular systems, HOMHs furnish a streamlined alternative, circumventing the complexities associated with synthesis procedures,²⁷ leading to the widespread application of various facile, low-temperature, and efficient solution-processing synthesis methods. Besides, the dimensionality of HOMHs is variable, which exerts a significant impact on their luminescent properties. Shrinking the dimensionality of HOMHs from three-dimensional (3D) to low-dimensional (2D, 1D,²⁸ 0D) can endow them with unique properties such as large exciton binding energy, multiple quantum wells and confinement effects,²⁹ controllable charge-carrier mobility, and efficient photoluminescence intensity.^{30–32} Low-dimensional metal halides have thus been witnessed to exhibit high photoluminescence quantum yield

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(PLQY) and broad emission range³³ on account of the formation of self-trapped excitons (STEs).³⁴ Organic cations play a key role in the formation of STEs and the distribution of hydrogen bond interactions within these HOMHs.³⁵ Recently, the integration of crown ethers into organic cations has been demonstrated with the exceptional ability to engage in intricate host-guest interactions with various metal cations and even organic ammonium cations.^{36–39} The introduction of crown ethers not only imparts a more diverse repertoire of hydrogen bonding interactions to the system, but also enhances the overall stability of the framework.⁴⁰ Additionally, the adaptable ring-opening diameter also offers the selectivity for distinct cations,⁴¹ enabling the modulation of luminescence in HOMHs.⁴² More importantly, the self-assembly systems of HOMHs, enabled by the presence of weak ionic bonds, low formation energies, flexible crystal nature, and multiple excited states, offer a handy and efficient avenue for facile modulation of the intrinsic optical characteristics exposed to external stimuli.^{23,26,43,44}

Chiral low-dimensional metal halides combine the merits of chiral materials and low-dimensional metal halides.⁴⁵ Chiral materials can manipulate the circular polarization states of light naturally.⁴⁶ Applying the strategy of chirality transfer from organic chiral ligands, the chiroptoelectrics of low-dimensional metal halides have been comprehensively explored and developed, including circularly polarized luminescence (CPL), which refers to the emission of light with a specific polarization state, wherein the electric field vector rotates in a circular manner.⁴⁷ Due to its distinctive optical properties,⁴⁸ CPL has found extensive applications⁴⁹ in crucial domains such as 3D imaging and printing, sensing and probing, anti-counterfeiting, quantum computing, *etc.*⁴⁹ Consequently, chiral low-dimensional metal halides pave a potential pathway for intelligent modulation of CPL by virtue of the integration of CPL emission and multi-stimuli responsiveness.²⁷

Herein, a pair of new zero-dimensional (0D) chiral antimony halides, *i.e.*, $[R/S\text{-}1,2\text{-propanediamine}(18\text{-crown-}6)_2]\text{SbCl}_6$, referred to as **R-Sb** and **S-Sb**, have been synthesized *via* antisolvent vapor-assisted crystallization. Within their crystalline configurations, the self-assembly behaviour of 18-crown-6 with chiral amine cations bestows an abundance of hydrogen bonding interactions and efficaciously segregates the inorganic octahedra, thereby boosting the photoluminescence efficiency (51.6% and 87.2% quantum yield for white and orange emission, respectively). Moreover, the chirality transfer engendered by the asymmetric hydrogen bonding interactions confers the system with effective CPL. More importantly, the chiral **R-Sb** and **S-Sb** exhibit multi-stimuli-responsive luminescence, showcasing their capacity to dynamically respond to excitation wavelength and temperature, enabling a back-and-forth emission colour switch from blue to orange emission. This study not only unveils a novel approach for the development of low-dimensional eco-friendly emitters with CPL, but also broadens the scope of candidate materials for optical and thermal sensing optoelectronic devices.

Results and discussion

Structural characterization

The target chiral 0D antimony halides (**R/S-Sb**) have been obtained *via* a facile antisolvent vapor-assisted crystallization (see details in the Experimental section). Their spatial configurations have been confirmed through single crystal X-ray diffraction (SCXRD) analyses. Chiral **R-Sb** and **S-Sb** crystallize in tetragonal chiral space groups of $P4_32_12$ and $P4_12_12$, respectively, subordinate to the 422 crystal class (Table S1†). In an asymmetric unit, a chiral 1,2-propanediamine cation (PDD²⁺) is anchored with double 18-crown-6 molecules through multiple N–H⋯OC hydrogen bonding interactions, featuring a host-guest architecture (Fig. 1a). The incorporation of the bulky 18-crown-6 isolates the inorganic $[\text{SbCl}_6]^{3-}$ octahedra effectively, forming the classical 0D configuration (Fig. 1b). Among the 0D metal halides, the large distances between $[\text{MX}_n]^{m-}$ units eliminate the electronic interactions and energy transfer between neighboring units, leading to a highly localized exciton in the excited state,^{50–53} facilitating the formation of highly emissive HOMHs. The steric effect of bulkier organic cations $[\text{PPD}(18\text{-crown-}6)_2]^{2+}$ enables the enhanced separation of $[\text{SbCl}_6]^{3-}$ octahedra, showing the shortest Sb⋯Sb distance of 10.11 Å for **R-Sb** and 9.66 Å for **rac-Sb** (Fig. S1a†). In comparison, in the absence of the crown, the longest distances between atoms of the chiral $[\text{S-PPD}]^{2+}$ cation along the crystallographic *b*- and *c*-axis directions are shortened to 5.26 Å and 3.42 Å (Fig. S2†), respectively. As a result, the shortest Sb⋯Sb distance of $[\text{S-}1,2\text{-propanediamine}]\text{SbCl}_6$ (**S-PPD-Sb**) shrinks to 6.22 Å. Different from the usual organic-inorganic hybrid antimony halides, the inorganic frameworks combine with organic cations through Sb–Cl⋯H–C hydrogen bonding interactions (Fig. S3†) rather than Sb–Cl⋯H–N interactions. By investigating the number of Sb–Cl⋯H–C hydrogen bonds concerning an inorganic

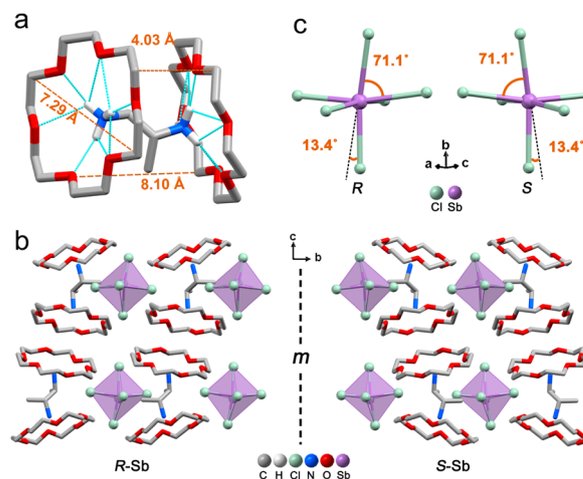


Fig. 1 Single-crystal structures of the chiral 0D antimony halides. (a) The structure of the $[\text{R-PPD}(18\text{-crown-}6)_2]^{2+}$ cation, showing host-guest interactions, size of 18-crown-6 (7.29 Å), longest distance (8.10 Å) and shortest distance (4.03 Å) between the two adjacent 18-crown-6 molecules. (b) Structural models of **R-Sb** and **S-Sb** (hydrogen atoms are omitted for clarity, *m* represents mirror plane). (c) Inorganic scaffold in **R/S-Sb**.



unit and the proportion of H \cdots Cl interactions around organic amines in **R-Sb** and **rac-Sb** structures by the Hirshfeld surface analysis method, it is revealed that **R-Sb** exhibits stronger hydrogen bonding interactions than **rac-Sb** (Fig. S4 and S5 \dagger).⁵⁴ Compared to the non-covalent interactions in **S-PPD-Sb**, the crown ether in the **R-Sb** structure obstructs the contact between the chiral amine cation and [SbCl₆]³⁻ octahedron, resulting in a decline in the proportion of H \cdots Cl interactions and the predominance of H \cdots O interactions (Fig. S6 \dagger). Moreover, the results of thermogravimetry-differential thermal analyses (TG-DTA) show that **R/S-Sb** preserves higher melting temperature than **rac-Sb** (Fig. S7 \dagger), consistent with the fact that the hydrogen bonding interactions are intensified in the chiral antimony chlorides. Under the effect of asymmetric hydrogen bonding interactions, severe octahedral distortion is observed in the chiral antimony chlorides (Fig. 1c). The hydrogen bonding with the axial chlorine atom induces a 13.4° tilt and the equatorial Cl-Sb-Cl bond angle was compressed to 71.1°. The corresponding bond angle variances (σ^2), as calculated *via* eqn (1):^{55,56}

$$\sigma^2 = \frac{1}{11} \sum_{i=1}^{12} (\theta_i - 90)^2 \quad (1)$$

reveal a greater extent of distortion of **R/S-Sb** compared to **rac-Sb**. The pronounced distortions in the octahedral geometry can be identified in **R-** and **S-Sb** with σ^2 of 71.0 deg.² significantly larger than that of **rac-Sb** (12.3 deg.²). The powder X-ray diffraction (PXRD) patterns of **R/S-Sb** as depicted in Fig. 2a accord well with the simulated results from the corresponding SCXRD data, verifying the high phase purity of the obtained single crystals.

Photophysical properties of chiral Sb halides

The obtained UV-vis absorption spectra reveal prominent absorption bands of **R/S-Sb** in the range of 200–400 nm,

accompanied by a minor absorption in the visible range (400–500 nm) (Fig. 2b), consistent with the pale-yellow colour observed in the crystals under ambient light. Due to the excited state splitting behaviours of the 5s² configuration like Sb³⁺, the two main absorption peaks around 213 nm and 315 nm can be attributed to characteristic ¹S₀ → ¹P₁ and ¹S₀ → ³P₁ electronic transitions of the Sb(III) ion containing 5s² lone-pair electrons.^{57,58} Under UV light irradiation at 365 nm, **R-Sb** exhibits intense orange emission (Fig. S8 \dagger). Therefore, steady-state photoluminescence (PL) and photoluminescence excitation (PLE) spectra (Fig. 2c) have been collected to investigate their luminescent behaviours. Upon excitation at the optimal excitation wavelength of 370 nm, **R/S-Sb** displays broadband emission centered at 612 nm, with a full width at half maximum (FWHM) of 116 nm (0.40 eV), and a significant Stokes shift of 242 nm (1.32 eV). Such a large Stokes shift signifies low self-absorption effects in the luminescent material, which is highly desirable for high-performance luminescent materials.^{29,59} Under the optimal excitation wavelength, **R-Sb** exhibits a high quantum yield of 87.2% (Fig. S9a \dagger). However, the **S-PDD-Sb** crystal is nonluminescent under ambient conditions (Fig. S10 \dagger), reflecting the key role of 18-crown-6 in the emission process. The luminescence lifetimes of **R-Sb** and **S-Sb** have been monitored by time-resolved decay PL spectra (Fig. 2d). The resulting decay curves could be well fitted with a single exponential function, with a lifetime of approximately 7.0 μs. Based on these aforementioned luminescence features of **R/S-Sb**, including a broad FWHM, large Stokes shift, and long luminescence lifetime (microsecond scale), it could be concluded that the emission of **R/S-Sb** centered at around 612 nm originates from triplet STE emission.^{60–62} The STE formation of **R/S-Sb** is contributed by its intrinsic low electronic dimensionality and soft lattice nature.⁶³

Stimuli-responsive luminescence of chiral Sb halides

For chiral **R-Sb** halides, a gradual blue-shift in the excitation wavelength engenders the gradual emergence of a high-energy emission peak at 492 nm. In stark contrast to the long-lived luminescence at 612 nm, which exhibits a lifetime of 7.0 μs, time-resolved PL decay spectroscopy reveals a significantly shorter lifetime of 9.3 ns at 492 nm (Fig. S11 \dagger), thereby signifying the single STE (¹P₁ → ¹S₀) electronic transition process.^{64,65} The time-resolved emission spectra of **R-Sb** with excitation at 365 nm were recorded to assign the dual emissions (Fig. 3c and d). The emission intensity at 492 nm rises steadily, reaching its maximum at 19.98 ns, before swiftly decaying and extinguishing within a short duration. Conversely, the emission peak at 612 nm exhibits a much slower decline, taking approximately 0.7 μs to diminish and eventually fade away. The distinct excitation spectra observed at 492 nm and 612 nm thus prove the involvement of different transition mechanisms for the dual emission bands (Fig. S12 \dagger). For the low-dimensional halides with the 5s²-configuration cations (such as Sb³⁺ or Sn²⁺), the *nsnp* excited states will be split into ¹P (singlet) and ³P (triplet) states under the impact of the Coulomb and exchange interactions.⁶⁶ Further, the ³P states will be split into non-

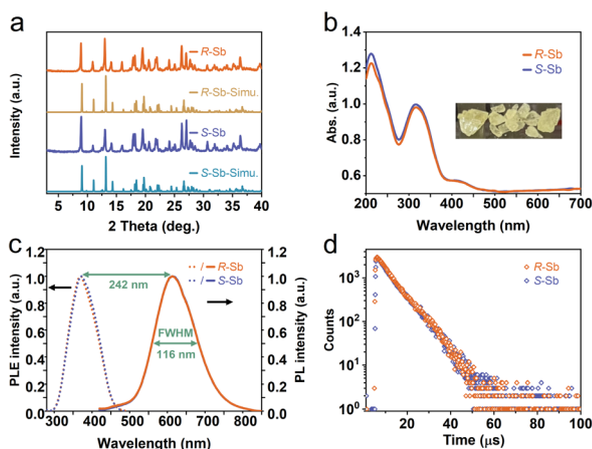


Fig. 2 Characterization of chiral OD antimony halides. (a) PXRD patterns and (b) UV-vis absorption spectra of **R/S-Sb**. Inset: optical image of **R-Sb** crystals under the radiation of natural light. (c) Normalized PL excitation (PLE), PL spectra and (d) time-resolved PL decay spectra of **R/S-Sb** at 612 nm.



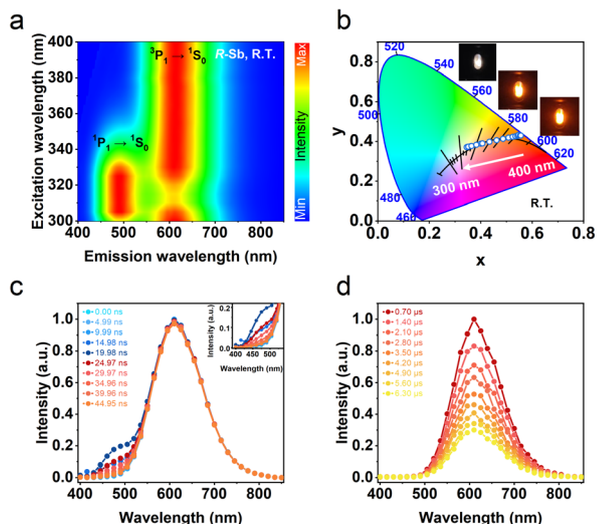


Fig. 3 Excitation wavelength-dependent PL properties of *R-Sb*. (a) Excitation–emission mapping patterns of *R-Sb* under ambient conditions. (b) The corresponding CIE chromaticity coordinates at different excitation wavelengths. Insets: luminescence photographs at excitation wavelengths of 300 nm (left), 360 nm (medium), and 380 nm (right). (c) and (d) Time-resolved emission spectra for *R-Sb* at the excitation wavelength of 365 nm. Note that $t = 0 \mu\text{s}$ ($t = 6.0 \mu\text{s}$) represents the start (end) of the excitation pulse and that the emission intensity reached its maximum at $t = 6.0 \mu\text{s}$.

degenerate states 3P_2 , 3P_1 , and 3P_0 by the spin–orbit coupling interaction (Fig. S13a†). The STE3 state should be converted from the STE¹ state by undergoing an intersystem crossing (ISC) process (Fig. S13b†), resulting to the delay emission process and long lifetime.⁶⁷

As the excitation wavelength transitions from 300 nm to longer wavelengths of up to 400 nm, a pronounced stimulus response is observed in the emission spectra of *R-Sb*. The emission at 492 nm is negligible for *R-Sb* when the excitation wavelength is longer than 350 nm, which can be ascribed to the direct $^3P_1 \rightarrow ^1S_0$ transition channel. The alteration of excitation wavelength can affect the exciton transition channel readily. Upon excitation at 320 nm, both singlet STE (STE¹) and triplet STE (STE³) states can be activated simultaneously, implying the energy transfer channel from STE¹ to STE³. Specifically, the high-energy emission intensity (I_{HE}) at 492 nm and the low-energy emission intensity (I_{LE}) at 612 nm initially increase before gradually diminishing upon the change of excitation wavelength from 300 to 400 nm, while the LE emission increasingly overshadows the one in the HE emission (Fig. 3a). This comprehensive variation in the relative intensities of the two peaks brings about a luminescence colour switch of *R-Sb* from the original yellowish white to yellow, and orange, resulting from the stimuli-responsive behaviours of excitation wavelength emission colours between STE¹ and STE³. Therefore, by controlling the excitation wavelengths, one could deftly generate the white emission (Fig. 3b). Blending a pair of complementary wavelengths, the CIE chromaticity coordinates of the white emission of *R-Sb* at 320 nm excitation were calculated to be (0.35, 0.37), obtaining high colour temperatures

(CCT) of 4109 K, along with a high PLQY of 51.6% (Fig. S9b†). Thus, the dynamic emission colour modulation can be realized facilely *via* targeted excitation of distinct STEs. For the multi-excitonic emission in a single system, the colour kinetic performance offers promising strategies for scheming multi-colour light-emitting diodes (LEDs) and multidimensional optical anti-counterfeiting materials.⁶⁸

To further investigate the photophysical mechanism of *R-Sb*, temperature-dependent PL spectra were recorded. Under excitation at a higher energy of 320 nm, the emission peak intensity at 492 nm gradually increases with lowering temperature (Fig. 4a). Furthermore, the fluorescence lifetime prolongs significantly from 5.2 ns at 320 K to 28.1 ns at 80 K (Fig. 4b), implying nonradiative recombination is suppressed in the cryogenic environment.⁶⁹ Additionally, the FWHM has noticeably narrowed down, indicating a weaker electron–phonon coupling interaction in the cryogenic environment.⁶⁰ Similar variations can be also observed at 612 nm in both luminescence intensity and lifetime (Fig. 4c and d). Under excitation by low-energy photons, the emission intensity at 492 nm becomes inconspicuous, signifying that only the STE³ state is actuated. In contrast, upon excitation by high-energy photons, the ground-state electrons experience excitation to higher energy states and are subsequently trapped by the STE¹ state. Then the electrons undergo an intersystem crossing (ISC) process, transiting from the STE¹ state to the STE³ state, and exhibiting dual-emission luminescence.

The dual-emission behaviour of *R-Sb* exhibits an exceptional temperature sensitivity, particularly in the vicinity of the 492 nm emission peak. Throughout the progressive cooling process from 320 to 80 K, the emission intensity at 492 nm experienced an obvious amplification by 17.3-fold, while the intensity at 612 nm saw a comparatively modest 2.4-fold increase. The temperature-dependent non-uniformity in the intensity variations of these two bands engenders a captivating colour-

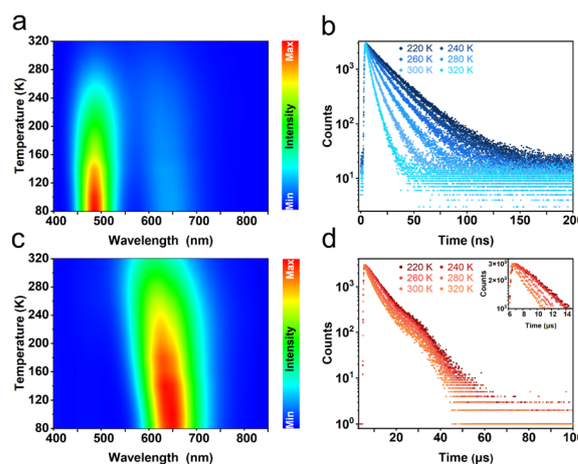


Fig. 4 Temperature-dependent PL measurements of *R-Sb*. (a) Temperature-dependent PL spectra ($\lambda_{\text{ex}} = 320 \text{ nm}$) and (b) decay PL spectra ($\lambda_{\text{ex}} = 300 \text{ nm}$) for *R-Sb*; (c) temperature-dependent PL spectra ($\lambda_{\text{ex}} = 370 \text{ nm}$) and (d) decay PL spectra ($\lambda_{\text{ex}} = 370 \text{ nm}$) for *R-Sb*. Inset: its partial amplification diagram from 6 to 14 μs .



switching response in the luminescence (Fig. S14a†). At 80 K, the prevailing emission dominance in the blue emission region, specifically at 492 nm, endows **R-Sb** with an overall blue-emitting appearance, as evidenced by the CIE coordinates of (0.18, 0.27). However, as the temperature gradually rises, the prominence of the blue emission wanes, prompting a transition in the overall luminescence to a white emission at 320 K, exemplified by the CIE coordinates of (0.35, 0.37).

The Huang–Rhys factor (S) is widely employed to evaluate the strength of electron–phonon coupling interactions in metal halides, which can be estimated *via* the following equation (eqn (2)).

$$\text{FWHM} = 2.36\sqrt{S}\hbar\omega_{\text{phonon}}\sqrt{\coth\frac{\hbar\omega_{\text{phonon}}}{2k_{\text{B}}T}} \quad (2)$$

where S refers to the Huang–Rhys factor, $\hbar\omega_{\text{phonon}}$ represents phonon frequency (\hbar is the reduced Planck's constant), and k_{B} is the Boltzmann constant. The function of FWHM (meV) to temperature (K) for HE and LE bands at the optimal excitation wavelength has been fitted and calculated. The fittings are exhibited in Fig. S15a–c†, where the S ($\hbar\omega_{\text{phonon}}$) values for HE and LE emission are 39.3 (18.6 eV) and 19.7 (26.2 eV), respectively. Based on the obtained values of $\hbar\omega_{\text{phonon}}$ for HE and LE, their self-trapping times (τ) were extracted using the expression $\tau = 2\pi/\omega_{\text{phonon}}$, and found to be $\tau_{\text{HE}} = 222.6$ fs and $\tau_{\text{LE}} = 157.8$ fs, respectively. The small τ values indicate the ultrafast formation of stable STE¹ and STE³ states.^{58,70} Large S factors for HE and LE bands imply strong electron–phonon coupling in the STE emission process on account of the large lattice distortion of chiral antimony halides.^{71,72} Moreover, according to the temperature-dependent PL spectra, the exciton binding energies for HE and LE emission can also be acquired by the following equation (eqn (3)), which describes the strength of the Coulomb attraction between an electron and a hole.^{73–75}

$$I(T) = I_0 / \left(1 + A \times e^{-\frac{E_{\text{b}}}{T \times k_{\text{B}}}} \right) \quad (3)$$

where E_{b} is the exciton binding energy, I_0 is the intensity at 0 K, and k_{B} is the Boltzmann constant. The E_{b} values of HE and LE emissions are estimated to be 121.2 meV and 202.6 meV, respectively. The two E_{b} values are both much higher than the thermal energy $T \times k_{\text{B}}$ at room temperature (26 meV), indicating that the stable STEs resist thermal decomposition.^{76,77} In addition, the smaller E_{b} of HE reflects the lower energy barrier for excitons to nonradiative centers. Therefore, the luminescence from the HE region is more susceptible to thermal quenching at room temperature.⁷⁸

CPL properties of chiral Sb halides

One of the fascinating optical characteristics exhibited by chiral metal halides is their response to and control of circularly polarized light. Through an analysis of the crystal structure presented earlier, it becomes evident that the chirality of organic cations can be effectively transferred to the inorganic framework through

asymmetrical hydrogen bonding interactions. This assertion finds validation in the circular dichroism (CD) spectra of the chiral antimony halides. The CD spectra of **R-** and **S-Sb** display mirror-image signals at 263 nm, 320 nm, and 356 nm, where all peaks are anterior to the absorption edge (Fig. 5a). This phenomenon is well known as the Cotton effect, associated with the splitting of electronic states under excitation in distinct CPL. The anisotropy factor, g_{CD} , is calculated *via* the following equation (eqn (4)):

$$g_{\text{CD}} = \frac{(A_{\text{L}} - A_{\text{R}})}{\text{absorption}} = \frac{\text{CD}[\text{mdegree}]}{32\,980 \times \text{absorption}[\text{o.d.}]} \quad (4)$$

where A_{L} and A_{R} are the absorption of left- and right-handed CPL. The corresponding pattern of absorption anisotropy factor g_{CD} is shown in Fig. 5c, which is up to 2.1×10^{-3} at 320 nm.

Given their chiral structures and strong photoluminescence properties, it is not surprising that **R-/S-Sb** displays pronounced CPL responses. The CPL signals of **R-** and **S-Sb** crystals manifest as a pair of mirror-symmetric curves in the range of 500–900 nm (Fig. 5b). Similarly, the extent of CPL is quantified and assessed by the luminescence anisotropy factor g_{lum} . The g_{lum} is defined as the difference in emission intensity of the material under excitation by two different circularly polarized lights, *i.e.*, $g_{\text{lum}} = 2(I_{\text{L}} - I_{\text{R}})/(I_{\text{L}} + I_{\text{R}})$,⁷⁹ where I_{L} and I_{R} are the intensities of left- and right-handed CPL. Fig. 5d implies the g_{lum} values of **R-** and **S-Sb** at the maximum emission wavelength are -5.7×10^{-3} and 5.2×10^{-3} , respectively, showing a high luminescence dissymmetry factor among the previously reported chiral antimony halides (Table S7†).

Density functional theory (DFT) calculations

To understand the PL nature of **R-Sb**, theoretical calculations based on the first-principles DFT have been performed. As depicted in Fig. 6a, the flat electronic bands indicate a localized electron band structure. Moreover, the band gap of **R-Sb** is direct, and the corresponding calculated PBE bandgap at M point is determined to be 3.67 eV, slightly lower than the experimental

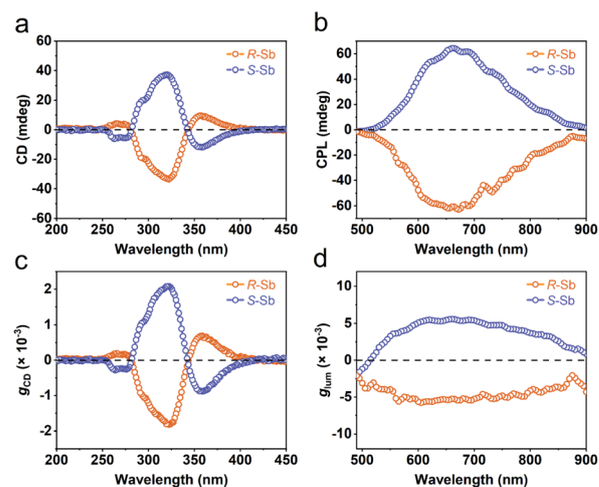


Fig. 5 Chiroptical properties of **R-/S-Sb**. (a) CD spectra, (b) CPL emission spectra, and the corresponding (c) g_{CD} factor and (d) g_{lum} factor of **R-/S-Sb**.



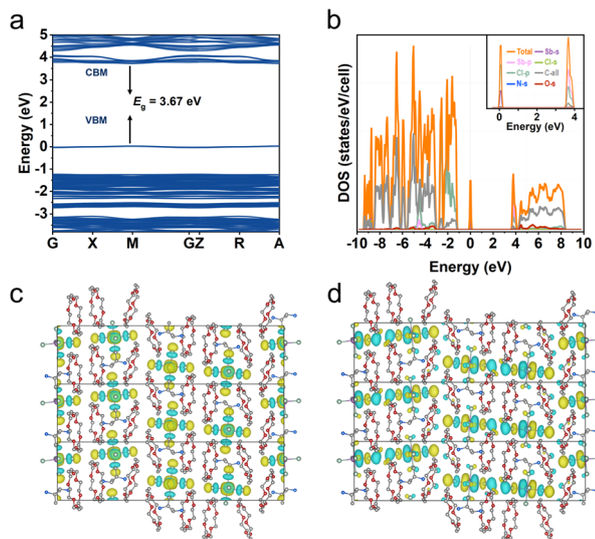


Fig. 6 DFT simulation of *R*-Sb. (a) Calculated band structure and (b) the corresponding PDOS for *R*-Sb. (c) The conduction band minimum (CBM) and (d) valence band maximum (VBM) of *R*-Sb.

value probably due to the discontinuity of the exchange-correlation function.⁸⁰ Both the conduction band minimum (CBM) and valence band maximum (VBM) are contributed from the localized electronic states of inorganic parts, which also agree with the calculated projected density of states (PDOS) (Fig. 6b). The CBM mainly consists of a mixture of Sb-p and Cl-p, while the VBM is mainly derived from Sb-s and Cl-p orbitals with dominant Cl-p orbitals. The electronic charge density patterns also unveil the electronic transfer from the halide to metal (Fig. 6c and d), implying luminescence process is primarily determined by the inorganic components.

Photophysical properties of achiral Sb halides

The UV absorption spectra of the achiral Sb halide *rac*-Sb indicate its similar absorption characteristics to those of *R*-Sb (Fig. S16[†]). The photophysical properties of *rac*-Sb are primarily assessed through its corresponding PL spectra. In the excitation-emission mapping patterns (Fig. S17a[†]), two emission centers can be observed, located at 490 nm and 626 nm, respectively. Time-resolved decay spectra reveal that the PL lifetimes at 490 nm and 626 nm are measured to be 7.2 ns and 7.4 μ s, respectively, suggesting that the luminescent behaviours at the two peaks can be attributed to the $^1P_1 \rightarrow ^1S_0$ and $^3P_1 \rightarrow ^1S_0$ transition processes, respectively. As shown in Fig. S17a,[†] switching the excitation wavelength from 300 nm to 400 nm results in non-synergistic alterations in the emission intensities of the two main peaks, with the highest PL intensity at 490 nm significantly higher than that at 626 nm. Similarly, as the excitation wavelength gradually redshifts, *rac*-Sb undergoes a significant emission transition from blue to white and eventually to orange (Fig. S17b[†]).

Nevertheless, the emission of *rac*-Sb is much lower than that of the chiral *R*-Sb. At the optimal excitation wavelengths of 319 nm and 371 nm, the PLQYs of *rac*-Sb are both below 15%.

This can be mainly attributed to the smaller degree of lattice distortion in the inorganic octahedra of *rac*-Sb, resulting in weaker electron-phonon coupling interactions. Temperature-dependent PL spectra measurements of *rac*-Sb at excitation wavelengths of 319 nm and 371 nm were conducted (Fig. S18[†]). By analyzing the relationship between the FWHM at 490 nm (HE) and 626 nm (LE) and temperature, the corresponding *S* of HE and LE can be determined to be 19.1 and 17.4, respectively (Fig. S19[†]), both lower than those of *R*-Sb, verifying the weaker electron-phonon coupling interactions in *rac*-Sb. Moreover, the denser arrangement of octahedra in *rac*-Sb also leads to the increased energy transfer between adjacent octahedra, thereby lowering the efficiency of photoluminescence.

Noticeably, *rac*-Sb can be readily prepared *via* the solvent-free synthesis. By mixing stoichiometric amounts of precursor materials in this ternary system and dry grinding the precursors in the glovebox, *rac*-Sb can be rapidly and facilely obtained. After grinding for 10 minutes, the samples can express the orange emission under the 365 nm UV radiation (Fig. S20b[†]). The PXRD pattern of the ground sample is fitted well with the simulated result from the SCXRD data (Fig. S20c[†]), testifying to the successful fabrication of *rac*-Sb. However, grinding the stoichiometric *R*-*S*-Sb precursor materials can only obtain the *rac*-Sb phase (Fig. S21[†]), probably due to the racemization of the chiral amine cation during the grinding process.

Conclusion

Adopting the host-guest self-assembly strategy mediated by a crown ether, we have designed and synthesized a pair of zero-dimensional chiral antimony halides (*R*-*S*-Sb). The inclusion of crown ethers not only creates rich hydrogen bonding networks, but also contributes to the formation of bulky chiral cations, segregating the $[SbCl_6]^{3-}$ inorganic octahedra and enhancing the PLQY, as high as 87.2% and 51.6% for orange and white emission, respectively. By virtue of the asymmetric hydrogen bonding interactions, the chirality of the organic cations is conveyed to the inorganic framework readily, inducing conspicuous asymmetrical lattice distortion within the octahedral structure. Consequently, *R*-*S*-Sb exhibits strong CPL emission, along with the good luminescence anisotropy factor ($|g_{lum}| \geq 5.2 \times 10^{-3}$), suggesting its potential as a circularly polarized LED (CP-LEDs). Intriguingly, the lattice distortion-induced dual-stimuli STE emission exhibits reversible emission transitions of chiral antimony halides from blue to white to orange colour, demonstrating sensitivity to both excitation wavelength and temperature. Meanwhile, the emission of *rac*-Sb also shares sensitivity to excitation wavelength, although it manifests a weaker emission than its chiral counterparts on account of the diminished electron-phonon coupling interactions. Moreover, *rac*-Sb can also be prepared readily *via* the solvent-free solid-state grinding pathway. This study provides a new protocol for the fabrication of a lead-free, efficient CPL emitter, while also expanding the application of chiral zero-dimensional metal halides in the field of multi-stimuli-responsive luminescence.



Experimental section

Materials

18-Crown-6 (99%) and *N,N*-dimethylformamide (DMF, 99.5%) were purchased from Aladdin. *R/S-rac*-1,2-propanediamine dihydrochloride (*R/S-rac*-PDD·2HCl, 98%) was purchased from Bidepharm. Antimony trichloride (SbCl₃, 99%) was purchased from Macklin. Tetrahydrofuran (THF, 99%) and dimethyl sulfoxide (DMSO, 99%) were purchased from Concord Technology (Tianjin). All reagents and solvents were used without further purification.

Synthetic procedures

Growth of *R/S-rac*-Sb single crystals. First, *R/S-rac*-PDD·2HCl (1.0 mmol, 0.113 g), 18-crown-6 (2.0 mmol, 0.534 g) and SbCl₃ (1.0 mmol, 0.230 g) were dissolved in a mixed solvent with 1.5 mL of DMF and 0.5 mL of DMSO at room temperature to form a clear precursor solution. The prepared precursor solution was filtered using a polyvinylidene fluoride (PVDF) filter with 0.50 μm pore-size to remove unreacted impurities. The filtrate was poured into a small vial (20 mL in volume) and then left open in a closed glass vial (300 mL) containing 30 mL of THF for 2–3 days. Pale-yellow transparent crystals of *R/S-rac*-Sb were filtered out and rinsed three times with THF.

Structural characterization

Single-crystal X-ray diffractions (SCXRD) were conducted on a Rigaku XtaLAB MM007 CCD diffractometer with Cu Kα radiation ($\lambda = 1.5418 \text{ \AA}$) or Mo Kα radiation ($\lambda = 0.71073 \text{ \AA}$) at 100 K. The structures were solved by SHELXT methods with the Olex2 programs and all non-hydrogen atoms were refined anisotropically by the least-squares technique on weighted F^2 using SHELXL. Powder X-ray diffraction (PXRD) spectra were recorded on a Rigaku D/Max-2500 diffractometer at 40 kV, 100 mA with a Cu-target tube and a graphite monochromator. Thermogravimetry analyses (TGA) were carried out using a standard TG-DTA analyser under an Ar flux of 10 mL min⁻¹ in the temperature range 30–800 °C at a heating rate of 10 °C min⁻¹.

Photophysical Characterization. UV-vis absorbance spectra were measured with a PerkinElmer LAMBDA 750 attached with an integrating sphere, which was calibrated by measuring BaSO₄ powder as a reference. Fourier transform infrared (FTIR) spectra of the compounds were collected on an FTIR spectrometer (SENSOR 37) from 4000 to 400 cm⁻¹. The excitation and emission of PL spectra at room temperature were conducted using an Edinburgh FS5 fluorescence spectrometer with a xenon lamp as the light source. The time-resolved PL decay spectra were collected with a microsecond flashlamp (μs scale) and TCSPC laser (ns scale). The temperature-dependent PL spectra (80–320 K) were measured on an Edinburgh FS5 equipped with a liquid nitrogen cooler and a linked temperature controlling stage. The PLQY was determined by a quantum yield measurement system attached to an integrating sphere with the optimal excitation wavelength. The transmission circular dichroism (CD) spectra of the prepared samples were measured and collected using a CD spectrometer (J-1500,

JASCO) with a scanning rate of 200 nm min⁻¹. The circularly polarized luminescence (CPL) spectra were recorded on a JASCO CPL-200 spectrometer.

Computational methods

DFT calculations were implemented in the DS-PAW adopting the projector augmented-wave method (PAW) to deal with the ion–electron interaction.^{81–83} We selected the generalized gradient approximation (GGA) of Perdew–Burke–Ernzerhof (PBE) to approximate the exchange–correlation energy.⁸⁴ Considering the nonbonding and long-range interaction in the hybrid organic–inorganic metal halides, the van der Waals (vdW) correction (DFT-D3) was adopted in the calculations.⁸⁵ The plane-wave basis set was defined by the energy cut-off at 450 eV. The energy convergence criterion for electronic iteration was set to be 10⁻⁵ eV. The structural relaxation was performed until the Hellmann–Feynman forces on each atom were less than 0.001 eV Å⁻¹. A *I*-centered 5 × 5 × 1 grid was adopted to simulate the *k*-space integrations.

Data availability

The data supporting this study are available within the main text and ESI.†

Author contributions

This article was written by X. H. after scientific discussion with all authors; J. X., B. X., and S. C. supervised the research; synthesis of single crystals and characterisations were carried out by X. H. with the help of S. H., Z. W., and J. G.; data treatments were performed by Y. Z., W. H., and R. S.; J. X. and Y. Z. revised the manuscript; theoretical calculations were performed by P. C.

Conflicts of interest

There are no conflicts to declare.

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