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Water-stable zero-dimensional hybrid zinc halide modulated by π – π interactions: efficient blue light emission and third-order nonlinear optical response†

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Achieving multifunctional optimization of halide optical materials through the precise modulation of intermolecular interactions is highly significant, yet it faces considerable challenges. Here, we propose a strategy of π – π interactions microregulating to achieve the simultaneous optimization of multiple properties of halide optical materials. Using this strategy, we obtained a new lead-free zero-dimensional (0D) zinc halide [DPE]ZnCl₄ (DPE = 1,2-di(4-pyridyl)ethylene), in which the protonated DPE cations are orderly arranged via π – π interactions, facilitating the ordered embedding and local regulation of [ZnCl₄]^{2–} units within long-range one-dimensional cationic π – π stacking. As a result of these modifications, [DPE]ZnCl₄ exhibits efficient blue light emission with a high photoluminescent quantum yield (PLQY) of 18.55%, far exceeding that of corresponding organic salt halides. Furthermore, this compound demonstrates an enhanced third-order nonlinear optical (NLO) response, with the modulation depth and the third-order NLO absorption coefficient reaching 0.70 and 3.81×10^{-10} m W^{–1}, respectively, surpassing those of three-dimensional (3D) perovskite quantum dots and most organic–inorganic hybrid halides. Notably, the modulation of π – π interactions results in a significant breakthrough in water resistance, allowing [DPE]ZnCl₄ to maintain excellent structural and performance stability in water for a week. This innovative strategy of π – π interaction modulation provides new avenues for the multifunctional regulation and water-proof design of halide optical materials, and it is expected to advance the development and functionalization of stable halide optical materials.

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Introduction

Three-dimensional (3D) lead halide perovskite quantum dots (PQDs) have several advantages, including a high light absorption coefficient, excellent photoelectric activity, efficient carrier transport and tunable emission wavelengths, making them promising for applications in light-emitting diodes (LEDs), solar cells, photodetectors, and X-ray scintillators.^{1–5} However, their low stability, high environmental sensitivity, lead toxicity, and aggregation-induced emission quenching present signifi-

cant challenges for optical applications.^{6–8} In this context, low-dimensional hybrid metal halides derived from 3D PQDs, featuring diverse structures and tunable emission properties, have emerged.^{9–11} Particularly for zero-dimensional (0D) hybrid metal halides, spatially isolated inorganic structural units are usually encapsulated by insulating organic matrices, forming a strong quantum confinement effect that promotes high carrier localization and enhances emission efficiency.^{12–16} By carefully selecting organic cations and metal halides, and controlling synthesis conditions, it is possible to achieve periodic embedding of halide units within organic cation assemblies, allowing for diverse structural and functional regulation. Consequently, combining environmentally friendly metal ions such as Sb³⁺, In³⁺, Mn²⁺, and Zn²⁺ with various types and functionalities of organic cations endows hybrid metal halides with enhanced prospects and significant future potential.^{17–21}

To date, a variety of strategies have been developed to effectively regulate the structure and photoluminescence (PL) properties of 0D hybrid metal halides, including adjusting the

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coordination configuration of metals, increasing distortion within the coordination configuration, optimizing the aggregation state of halide units, and expanding the distance between isolated units.^{22–26} Numerous 0D lead-free metal hybrid halides with outstanding performance have been reported and widely applied in LEDs, X-ray scintillation, anti-counterfeiting, and information encryption–decryption.^{27–30} However, the design and performance optimization of most halides are primarily achieved through the modification and regulation of anionic groups. In contrast, the synthesis and functional regulation of 0D halides guided from the perspective of organic cations remain relatively rare. Significantly, as an indispensable component of hybrid metal halides, organic cations play a role beyond merely charge balance and the confinement and regulation of halide units.^{31,32} Their intrinsic properties and potential functions warrant further exploration. In particular, designing and regulating the packing modes and intermolecular interactions between organic cations and metal halides is expected to further manipulate their photophysical properties and even introduce additional functional attributes, achieving multifunctionalization of new hybrid metal halide materials.^{33–35} For instance, Lei *et al.* used [BTPP]⁺ (BTPP = benzyltriphenylphosphonium) as an organic cation to obtain highly efficient dual-emission zinc-based halide phosphorescent materials through the heavy atom effect.³⁶ In this material, the ZnBr₄^{2–} unit exhibits strong blue emission derived from self-trapping exciton (STE), while the [BTPP]⁺ cation shows efficient green phosphorescence emission. This synergistic regulation of organic–inorganic components significantly activated the originally dormant phosphorescence properties of organic cation, resulting in highly efficient and long-lifetime room temperature afterglow, which utilized in time-color dual-resolved optical anti-counterfeiting and information encryption. Therefore, regulating the optical activity and functionality of metal halides through organic components and their spatial interactions is of great significance and presents considerable challenges.

Except for photophysical activity and functionalization, the extreme sensitivity and instability of hybrid metal halides to water and humid air further exacerbate their predicament.^{37–40} This weak resistance to water and humidity not only diminishes the intrinsic activity and application potential of these materials but also poses serious environmental challenges due to the release of metal ions following hydrolysis, particularly Pb²⁺.^{33,41,42} Therefore, improving water stability is essential for advancing the development of perovskites and hybrid metal halides. Although the stability of PQDs has been significantly enhanced through porous material encapsulation strategies, the complexity of these processes and the risks of leakage after encapsulation have prompted researchers to seek more effective strategies.^{43–48} Consequently, developing new methods to improve the inherent water stability of metal halides is urgently needed.

Inspired by the design concepts of metal–organic frameworks (MOFs) and hydrogen-bond organic frameworks (HOFs), introducing strong conjugated groups to form robust supramo-

lecular structures connected by strong non-covalent interactions is expected to enhance the water resistance of the system.^{49–51} These strong non-covalent interactions between conjugated groups offer greater stability than hydrogen bonds with water molecules, effectively preventing water erosion of the halide units protected by organic cations and thus avoiding metal ion hydrolysis. Additionally, these supramolecular structures connected by non-covalent interactions could also promote rapid charge transfer, leading to new functionalities inspired by ultrafast charge transport.⁵² Thus, designing and screening specific organic cations and rationally manipulating their spatial interactions are expected to create highly stable and multifunctional halide optical materials, offering new prospects for the revival of metal halides.

Based on the above considerations, here we selected 1,2-di(4-pyridyl)ethylene (DPE) with bidirectional conjugated structures as the organic cation and template agent, and synthesized a hybrid lead-free metal halide [DPE]ZnCl₄ based on isolated [ZnCl₄]^{2–} unit. As expected, the protonated DPE cations are arranged orderly through π – π interactions, forming a one-dimensional (1D) supramolecular stacking. This regulation of intermolecular non-covalent interactions not only significantly improves blue light emission efficiency but also imparts the material enhanced third-order nonlinear optical (NLO) activity, with the modulation depth and third-order NLO absorption coefficients surpassing those of 3D PQDs and most organic–inorganic hybrid halides. Most importantly, modulation of π – π interactions has led to a significant breakthrough in water resistance of halides maintaining excellent structural and performance stability after immersion in water for a week. This π – π interaction regulation strategy involving conjugated organic cations paves the new way for the precise design and long-term utilization of stable and multifunctional hybrid metal halide optical materials.

Results and discussion

Using DPE as the organic building block and ZnCl₂ as the metal source, colorless crystals of [DPE]ZnCl₄ were obtained through a self-assembly reaction under solvothermal conditions. X-ray single crystal diffraction analysis reveals that [DPE]ZnCl₄ crystallizes in the monoclinic space group *C2/c*, containing isolated [ZnCl₄]^{2–} tetrahedral anions, protonated DPE cations and solvent water molecules (Fig. 1). In the [ZnCl₄]^{2–} anion, each Zn²⁺ ion is coordinated with four Cl[–], forming a slightly distorted tetrahedral structure, with the Zn–Cl bond lengths ranging from 2.26 to 2.28 Å. Interestingly, the protonated [DPE]²⁺ cations are arranged orderly through strong π – π interactions from pyridine rings with adjacent cations, with a distance of 3.67 Å between the central rings of adjacent cations. This arrangement results in long-range 1D cationic π – π stacking, which contribute to enhancing the structural stability for [DPE]ZnCl₄ and the interaction and energy transfer of localized electrons. All [ZnCl₄]^{2–} polyhedra are periodically embedded into the DPE organic matrix, forming a

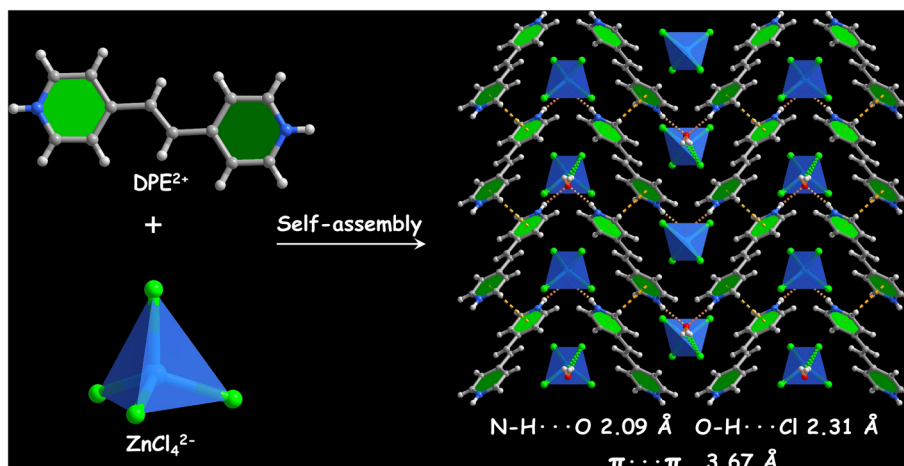


Fig. 1 Schematic diagram of structure assembly of [DPE]ZnCl₄. Color legends: cyan, Zn; green, Cl; blue, N; gray, C; red, O; white H.

perfect 0D hybrid structure. In this structure, the hydrogen on the pyridine N forms strong hydrogen bonds (N-H...O 2.09 Å) with free water molecules, while the hydrogen on the water molecules forms weak hydrogen bonds (O-H...Cl 2.31 Å) with the Cl⁻ from [ZnCl₄]²⁻. These multiple hydrogen bonds and strong π - π interactions ensure the firm integration of the organic cation [DPE]²⁺ and the inorganic anion [ZnCl₄]²⁻ within the hybrid structure. The experimental powder X-ray diffraction (PXRD) patterns of [DPE]ZnCl₄ matched well with the simulated based on single-crystal structural, indicating a phase-pure sample (Fig. S1†). The thermal stability of [DPE]ZnCl₄ was characterized by thermal analysis, and the results show that the compound can remain stable up to 220 °C (Fig. S2†). The morphology of [DPE]ZnCl₄ was characterized using field emission scanning electron microscopy (FE-SEM), revealing a ribbon-like structure with a length exceeding 100 μ m (Fig. S3†). Energy dispersive spectroscopy (EDS) and elemental mapping images indicated that Zn, Cl, C, and N are uniformly distributed throughout the crystal (Fig. S4†).

The optical properties of [DPE]ZnCl₄ were characterized by solid-state UV-Visible absorption, steady-state, and time-resolved PL emission spectroscopy. The UV-Vis absorption spectrum indicates a strong absorption peak around 305 nm with the cutoff wavelength at 376 nm (Fig. S5a†). The band gap of [DPE]ZnCl₄ was estimated to be 3.27 eV according to the Tauc formula, which corresponds to the colorless appearance (Fig. S5b†). As shown in Fig. 2a, under the irradiation of 365 nm ultraviolet light, the colorless needle-shaped crystals of [DPE]ZnCl₄ emit a strong blue light. The compound exhibits a broad excitation spectrum in the 200–400 nm range, accompanied by two excitation peaks at 282 nm and 370 nm. Under ultraviolet excitation at 370 nm, [DPE]ZnCl₄ exhibits a narrowband blue emission with a maximum emission peak at 429 nm (Fig. 2b). The full width at half maximum (FWHM) and the Stokes shift were measured to be 29 and 59 nm, respectively, which are smaller than that of reported hybrid

zinc halides, such as (EP)ZnBr₄ (79 nm), (TMPDA)ZnBr₄ (78 nm), [H₂MPz]ZnBr₄ (86 nm).⁵³ The narrowband emission has the Commission Internationale de l'Eclairage (CIE) chromaticity coordinates of (0.174, 0.072), corresponding to the observed blue emission (Fig. 2c). Under the ultraviolet light excitation at 370 nm, the PLQY of [DPE]ZnCl₄ was measured as 18.55%, significantly exceeding that of the inorganic hybrid Cs₂ZnBr₄ (< 5%) and many other reported blue-light emitting perovskite halides (Fig. S6†).^{54–56} To elucidate the dynamics of its radiative transitions, time-resolved PL spectra were measured at room temperature with the maximum emission wavelength of 429 nm. As shown in Fig. 2d, the PL decay curve can be fitted with a single exponential function to give a short lifetime of 8.12 ns, which indicates a single radiation path. Considering its broad excitation spectrum, we further conducted emission spectra related to excitation wavelengths at 300 K. Under ultraviolet excitation in the range of 230–400 nm, [DPE]ZnCl₄ exhibits the same narrowband emission profile and only one dominant emission center in the 3D excitation and emission correlation map, indicating a single radiative PL mechanism (Fig. 2e and Fig. S7†). To further understand the PL mechanism, we analyzed the PL spectrum of organic cation salts of [DPE]Cl₂, which exhibited a similar emission spectrum to [DPE]ZnCl₄ with the maximum emission peak at 405 nm upon exciting with 329 nm light (Fig. S8†). The similar emission spectrum shape and emission position indicate that the PL of [DPE]ZnCl₄ originates from organic salts. Moreover, the time-resolved PL spectrum of the organic salt can be fitted by a single exponential function with a lifetime of 2.66 ns, comparable to that of [DPE]ZnCl₄, further elucidates that the emission of [DPE]ZnCl₄ is attributed to the organic salt [DPE]Cl (Fig. S9†). It is worth noting that compared with the high PLQY of [DPE]ZnCl₄, the organic salt emitted almost no light under ultraviolet light, accompanied by negligible PLQY. This indicates that the filling of halide anions and the ordered π - π interaction of molecules effectively suppress the fluorescence quenching effect and the probability

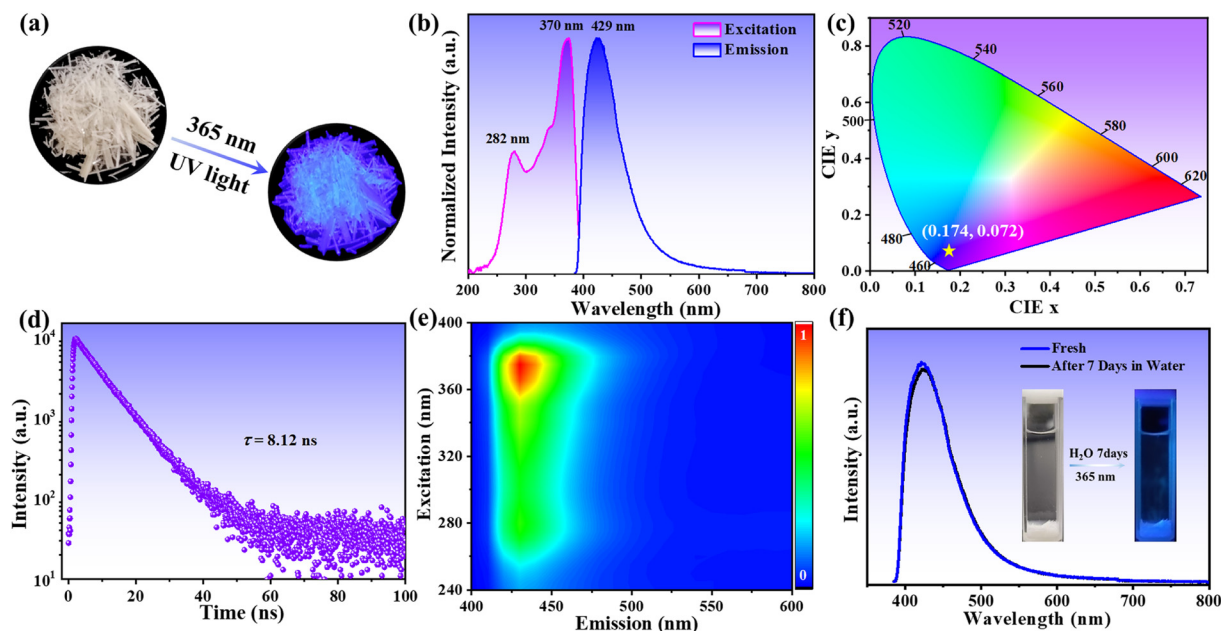


Fig. 2 Photoluminescence characterizations of [DPE]ZnCl₄. (a) Photos of crystals in sunlight and ultraviolet light of 365 nm; (b) excitation and emission spectra; (c) the CIE coordinates; (d) the PL decay curve at 429 nm at 300K; (e) the 3D excitation and emission correlation map; (f) comparison of the PL spectrum before and after immersion in water for 7 days.

of non-radiative transition, thereby significantly enhancing its PL efficiency.

Due to the strong π - π interactions between molecules, we further investigated the water stability of [DPE]ZnCl₄ by immersing the crystals in distilled water for a week. The results demonstrated that after one week of immersion, the crystals maintained both their shape and transparency, while continuing to emit bright blue light under ultraviolet light (Fig. 2f). The PL spectra of freshly prepared samples and those immersed in water for one week remained consistent, with no significant decrease observed, thereby indicating remarkable stability. Similarly, PXRD spectrum of samples after one week of immersion showed that all peaks matched well with those of the fresh samples, confirming their structural stability (Fig. S10[†]). In addition, the morphology of [DPE]ZnCl₄ showed no significant change before and after immersion in water for 7 days (Fig. S11). To determine whether the samples dissolved in water, we evaporated the separated water and found no obvious precipitate after complete evaporation. Additionally, inductively coupled plasma mass spectrometry (ICP-MS) testing of the solution after immersion showed a Zn²⁺ concentration of 3.2 mg L⁻¹, further indicating the negligible leaching of Zn ions from the material. These results demonstrate that by regulating the π - π interactions between organic cations, the water stability of the halide can be significantly enhanced, providing an effective solution for reducing environmental pollution and enabling its application under harsh conditions.

The strong π - π interactions in [DPE]ZnCl₄ prompted further investigation into its third-order NLO properties. We combined [DPE]ZnCl₄ with polydimethylsiloxane (PDMS) to prepare a flexible, transparent [DPE]ZnCl₄@PDMS film. Under sunlight,

this composite film exhibits high transmittance, allowing clear recognition of patterns behind the film (Fig. 3a). Under ultraviolet light irradiation, the film displays uniform bright blue luminescence, indicating that [DPE]ZnCl₄ is uniformly distributed within the polymer matrix (Fig. 3b). Notably, this film demonstrates good flexibility, allowing continuous folding and stretching without altering its shape, making it suitable as a flexible PL and third-order NLO material. The third-order NLO properties were evaluated using a standard open-aperture Z-scan system with a 532 nm nanosecond laser and a pulse energy of 4 μ J. At a laser pulse energy of 90 μ J, the pure PDMS film did not exhibit any nonlinear optical response, thus ruling out the influence of the substrate. As shown in Fig. 3c, the organic salt [DPE]Cl₂ exhibited slight reverse saturable absorption with a minimum normalized transmittance of 0.95 at its focus ($Z = 0$), demonstrating a weak third-order NLO response. In comparison, [DPE]ZnCl₄ exhibited an enhanced third-order nonlinear response with the modulation depth of up to 0.70, indicating excellent optical limiting properties. Further quantitative assessment of their third-order NLO responses, obtained by fitting the Z-scan curves, yielded a third-order nonlinear coefficient (β) for [DPE]ZnCl₄ of 3.81×10^{-10} m W⁻¹, which is 7.3 times greater than that of the organic salt [DPE]Cl₂. Additionally, the optical limiting thresholds (F_{OL} , defined as the input fluence at which the transmittance is half of the linear transmittance) for [DPE]Cl₂ and [DPE]ZnCl₄ were calculated to be 3.45 and 2.23 J cm⁻², respectively (Fig. 3d). These results indicate that by employing the π - π interactions microregulation strategy, we achieved a significant enhancement in the third-order nonlinear coefficient and modulation depth of [DPE]ZnCl₄, surpassing those

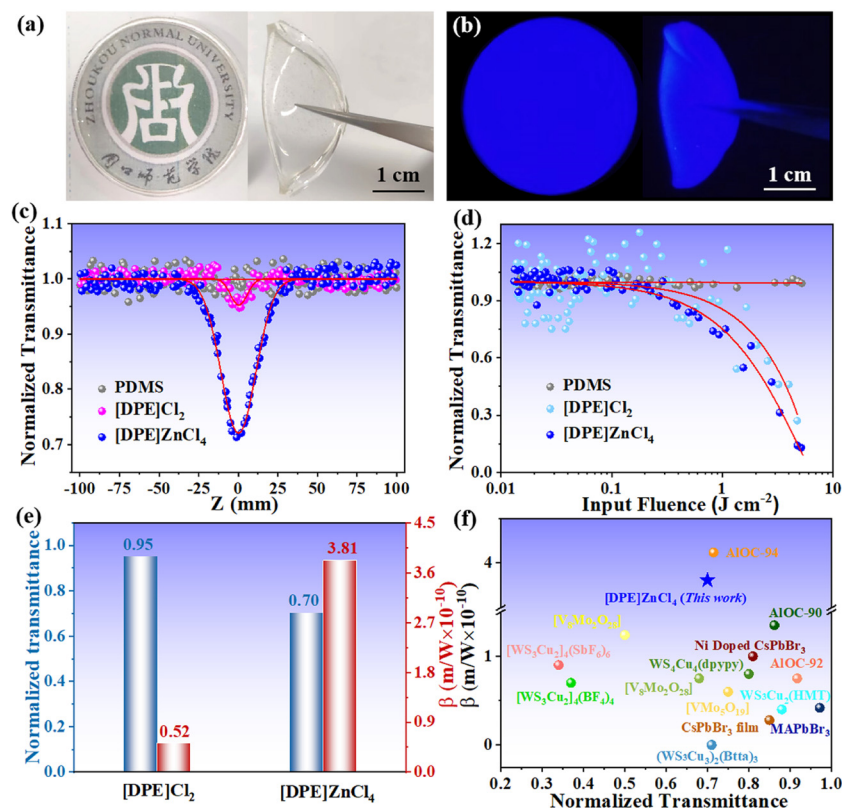


Fig. 3 (a and b) Photos of [DPE]ZnCl₄@PDMS films in sunlight and ultraviolet light of 365 nm; (c) the open-aperture Z-scan diagrams of [DPE]Cl₂ and [DPE]ZnCl₄ at 532 nm; (d) the curves of normalized transmittance versus input fluence; (e and f) comparison of the minimum normalized transmittance and the nonlinear absorption coefficients for [DPE]Cl₂, [DPE]ZnCl₄ and some representative crystalline materials reported.

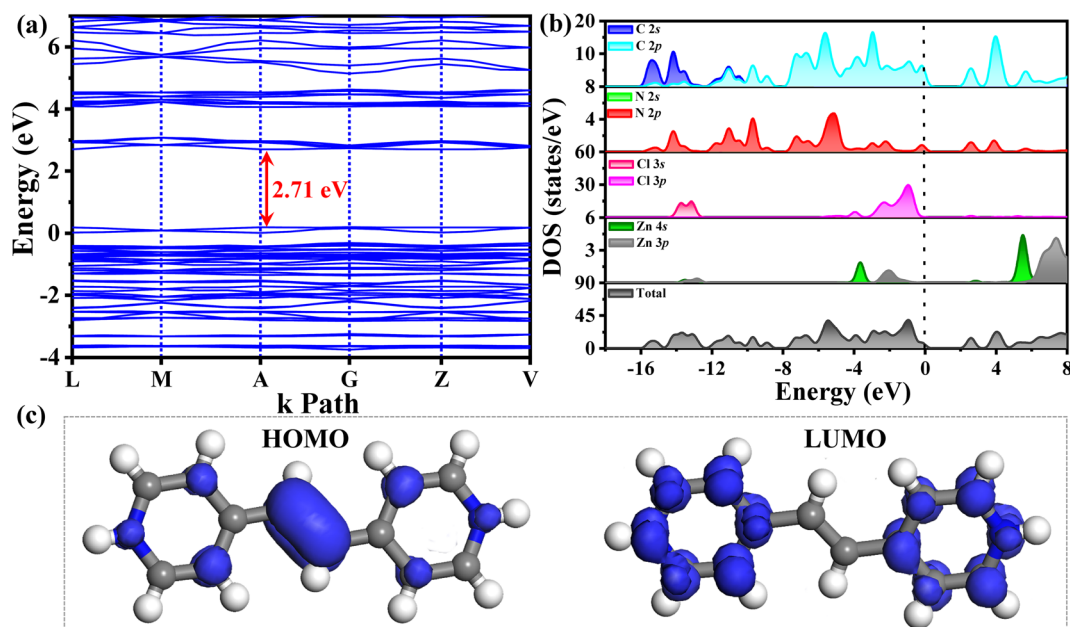


Fig. 4 Band structures and density of states of [DPE]ZnCl₄ (a and b); The frontier molecular orbitals (HOMO and LUMO) of [DPE]ZnCl₄ (c).

of the original organic salts, most PQDs materials, and other crystalline materials (Fig. 3e, f and Table S1†). Further precise structural analysis indicates that compared to the π - π interactions ($d_{\text{Cg-Cg}} = 4.75 \text{ \AA}$) in the organic salt of [DPE]Cl, the incorporation and reassembly of the $[\text{ZnCl}_4]^{2-}$ structural unit significantly enhance the π - π interactions ($d_{\text{Cg-Cg}} = 3.67 \text{ \AA}$) among the organic cations in [DPE]ZnCl₄ (Fig. S12†). This enhancement of intermolecular interactions leads to a significant improvement in the modulation depth and the NLO coefficients. This simple and effective microregulation strategy provides a new approach for developing high-performance hybrid metal halide third-order NLO materials.

Theoretical calculations based on density functional theory (DFT) were conducted to analyze the electronic structure and band gap of [DPE]ZnCl₄. As shown in Fig. 4a, the band structure calculations indicate that [DPE]ZnCl₄ has a direct band gap of 2.71 eV, consistent with the experimental band gap obtained from solid-state UV-Vis measurements. Furthermore, density of states (DOS) analysis shows that the conduction band minimum (CBM) and the valence band maximum (VBM) are primarily derived from the C-2p and N-2p states in the DPE cation, indicating that the organic cations significantly contribute to the band structure and electronic properties of [DPE]ZnCl₄ (Fig. 4b). This further supports the conclusion that PL originates from the organic cations. Orbital distribution analysis confirms that the lowest unoccupied molecular orbital (LUMO) is primarily located on the C=C bonds of the organic cation, while the highest occupied molecular orbital (HOMO) is chiefly situated on the pyridine ring, facilitating rapid ligand-to-ligand charge transfer (LLCT) (Fig. 4c). Consequently, strong intermolecular π - π interactions link organic cations to form non-covalently linked molecular wires, further enhancing efficient charge transfer within the lattice and boosting the third-order NLO properties.

Conclusions

In summary, we have developed a microregulation strategy involving π - π interactions to construct highly water-resistant multifunctional zinc halide optical materials. As a typical example guided by this strategy, [DPE]ZnCl₄ exhibits an ordered one-dimensional cation stacking mode induced by π - π interactions, thereby enhancing the rigidity and stability of the structure, effectively facilitating the regulation and efficient transfer of localized charges. With the microregulation of the aforementioned π - π interactions, [DPE]ZnCl₄ demonstrates efficient blue light emission and enhanced third-order NLO activity, with its PLQY and NLO coefficient significantly surpassing those of 3D PQDs and most organic-inorganic hybrid halides. More importantly, this simple structural microregulation strategy enhances the water resistance of [DPE]ZnCl₄, which can be stable in water for more than 7 days. This work realizes synergistic regulation and simultaneous optimization of multiple optical functions and stability in metal halides, providing new insights into the design and application of

novel multifunctional metal halide perovskite materials with high stability, potentially expanding their applications across a broad chemical range and under extreme conditions.

Data availability

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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