

Role of Metal–Chloride Anions in Photoluminescence Regulations for Hybrid Metal Halides

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ABSTRACT: Organic–inorganic hybrid metal halides with emissive organic cations are of great interest due to their structural diversity and interesting photophysical properties. Here, we assemble emissive organic cations (EnrofloH₂²⁺) with different metal–chloride anions (Pb₂Cl₆²⁻ to Bi₂Cl₁₀⁴⁻ to SnCl₆²⁻) to form the new single crystal phases, and thus the photoluminescence properties of the metal halides, including Stokes shift, full width at half-maximum (FWHM), and photoluminescence quantum yield (PLQY) have been studied accordingly. (EnrofloH₂)SnCl₆·H₂O, as an example, possesses narrow FWHM and high PLQY, which are caused by the strong π – π stacking and inter- and intramolecular hydrogen bonds interactions. Compared with EnrofloH₂²⁺ cation in solution, the interactions generate a restraining effect and increase the rigid degree of EnrofloH₂²⁺ cation in the bulk single crystals. Our work clarifies the photophysical properties of the EnrofloH₂²⁺ organic cations by constructing the inter- and intramolecular interactions and boosts the further study of organic–inorganic hybrid metal halides materials with different luminescence mechanisms.



here has been a great development of organic-inorganic hybrid metal halides due to their unique advantages, such as solution-process, low cost, large-area production, and various applications.¹⁻³ The flexible selectivity of organic cations and center metal ions (Cu⁺, Pb²⁺, Sn²⁺, Ge²⁺, Bi³⁺, Sn⁴⁺, etc.) in these compounds allows a wide choice of the compositions for elements in achieving the desired properties for photonic applications, such as photovoltaic cells, photodetectors and light-emitting diodes (LEDs).⁴⁻⁷ In general, the organic cations are divided into emissive organic cations and nonemissive organic cations according to the effect of organic cations on the photoluminescence (PL) behaviors of hybrid metal halides. The nonemissive organic cations only have a contribution to the construction of crystal structure for metal halides. In contrast, the emissive organic cations not only play a role in constructing the crystal structure but also have a contribution to the PL. Generally, luminescent metal halides exhibit free excitons (FEs) emission and self-trapped excitons (STEs) emission, related to the bandgap and the degree of lattice distortion for metal-halogen anions, which are found both in nonemissive and in emissive organic cation-containing compounds.⁸⁻¹⁰ Especially, metal halides with emissive organic cations have flexible selectivity, and their PL comes from either the individual organic cations or the combination of the organic cations and STEs.¹¹⁻¹³

Moreover, the emission of individual organic cations can be also regulated by selecting different metal—halogen anions.^{14,15} As a result, reasonably assembling organic cations with metal halogen anions to form the single crystals can alter the luminescence properties of hybrid metal halide materials for future burgeoning optoelectronic applications.

Enrofloxacin, as an emissive organic molecule, is composed of the piperazinyl and quinoline groups (Figure 1). They can



Figure 1. Chemical structure of Enrofloxacin (Enroflo).

be assembled with metal-halogen anions to form both complexes and ionic compounds.¹⁶ Furthermore, the Enrofloxacin molecule can exist as cationic, zwitterionic, and anionic species in solution on the basis of different pH conditions, whose emissions are closely related to the degree of protonation.^{17–20} In addition, their PL also can be controlled

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Figure 2. Calculated and experimental X-ray powder patterns of ENR-Pb (a), ENR-Bi (b), and ENR-Sn (c) crystals. Optical microscopic images of ENR-Pb (d, g), ENR-Bi (e, h), and ENR-Sn (f, i) single crystals. (d)–(f) daylight; (g)–(i) 365 nm UV light.

by regulating their local environment. It is well-known that the PL of pure organic cations are closely related to their rigidity. The bulk assembly of emissive organic cations and metalhalogen anions can form a hybrid metal halide, which can provide the rigid frame caused by hydrogen bonds and $\pi - \pi$ interactions. Furthermore, in compounds, the flexibility of pure organic cations is decreased; thus, the tunable PL for pure organic cations is realized.²¹⁻²⁶ And thus, several functionalized metal halides are obtained by assembling pure organic cations with multifarious metal-halogen anions. Here, we reported three new hybrid metal halides, (EnrofloH2)Pb2Cl6. H_2O (abbreviated as ENR-Pb), (Enroflo H_2)BiCl₅·Cl·2(H_2O)· H₃O (abbreviated as ENR-Bi), and (EnrofloH₂)SnCl₆·H₂O (abbreviated as ENR-Sn). The theoretical calculation and experimental results indicated that the emissions of these three single crystal compounds come from the singlet state fluorescence emissions of $EnrofloH_2^{2+}$ organic cations. Interestingly, the fluorescence emissions of $EnrofloH_2^{2+}$ in these single crystal compounds have large differences, such as maximum emission wavelengths, full width at half-maximum (FWHM), Stokes shift, and PLQY due to the discrepant confined environment of the organic cations. In as-synthesized metal halides, the organic cations and metal-chloride anions are connected by hydrogen bonds, and the hydrogen-bond networks in the environment of organic cations enables enhancement of their rigid degree. Moreover, serried packing of EnrofloH22+ in different hybrid metal halide can provide

stronger $\pi - \pi$ stacking interactions, which further improve the rigid degree of organic cations.²⁷ It is worth noting that there are both inter and intramolecular hydrogen bonds and $\pi - \pi$ interactions in ENR-Sn, the emission spectrum has the smallest FWHM with 40 nm and Stokes shift with 30 nm, and the PLQY of up to 19.23 %. As a result, organic–inorganic hybrid metal halides synthesized by emissive organic cations show better optical properties than single organic cations. Our work not only provides a series of new hybrid metal halide materials for future emerging optoelectronic applications but also provides an essential way to adjust the optical properties of the pure organic cations by rational assembly.

The crystallographic information files (CIF) of the assynthesized single crystal compounds, ENR-Pb, ENR-Bi, and ENR-Sn, are demonstrated in the Supporting Information, and the main crystal data are shown in Table S1. The comparisons of experimental and calculated X-ray diffraction (XRD) patterns of ENR-Pb, ENR-Bi, and ENR-Sn single crystal compounds are demonstrated in Figure 2a-c. It is worth mentioning that there are some differences between the calculated and experimental XRD patterns due to the preferred crystallite orientation. The optical microscopic images of these as-synthesized metal halide single crystals under daylight and ultraviolet light (UV) are described in Figure 2d-i. All of the single crystals show the yellow colors in daylight. Under the excitation of 365 nm UV light, these three single crystals show the bright blue fluorescence. The ENR-Pb single crystal

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Figure 3. Crystal structure diagrams and asymmetric parts of unit cell structures of ENR-Pb (a, a'), ENR-Bi (b, b'), and ENR-Sn (c, c') single crystals: $PbCl_6^{4-}$, green; $BiCl_6^{3-}$, cyan; $SnCl_6^{2-}$, yellow. Atoms: Pb, green; Bi, cyan; Sn, yellow; N, blue; O, red; F, orange; C, gray; H, white, and the color of Cl atoms are in agreement with the corresponding colors of metal-chloride anions.



Figure 4. Normalized PL and PLE spectra of ENR-Pb (a), ENR-Bi (b), and ENR-Sn (c) single crystals at room temperature. The arrows in (a)-(c) show the Stokes shifts. Decay curves of ENR-Pb (d), ENR-Bi (e), and ENR-Sn (f) single crystals at room temperature.

crystallizes in centrosymmetric monoclinic space group $P2_1/c$. The asymmetric part contains one ion of $C_{19}H_{24}FN_3O_3^{2+}$ (Enroflo H_2^{2+}), two Pb²⁺ ions, six coordinated Cl⁻ ions, and one H₂O molecule (Figure 3a'). The main bond lengths are shown in Tables S2. Each Pb²⁺ ion is coordinated by six Cl⁻ ions and forms PbCl₆⁴⁻ metal-chloride anions. It is of interest to note that the PbCl₆⁴⁻ metal-chloride anions. It is of interest to note that the PbCl₆⁴⁻ are connected by nodes-sharing, edge-sharing, and face-sharing, forming infinite double chain (Figure 3a). There are O-H…Cl, O-H…O, N-H…Cl, C-H…Cl, and C-H…O hydrogen bonds in the structure (Table S3), three of which are intramolecular hydrogen bonds. Intermolecular

hydrogen bonds form 3D net, which joints the Enroflo H_2^{2+} cations with PbCl₆⁴⁻ metal-chloride anions chains and H₂O molecules. The ENR-Bi single crystal crystallizes in triclinic space group $P\overline{1}$. The asymmetric part contains one $C_{19}H_{24}FN_3O_3^{2+}$ (Enroflo H_2^{2+}), one Bi³⁺ with five coordinated Cl⁻ ions and one Cl⁻ isolated ion, two H₂O molecules, and one H₃O⁺ ion (Figure 3b'), so that the total charge of the unit cell is zero. The main bond lengths are shown in Tables S4. The Bi³⁺ ion is coordinated by six Cl⁻ ions and forms BiCl₆³⁻ metal-chloride anions. The adjacent two octahedrons are linked with each other by edges forming a OD block Bi₂Cl₁₀⁴⁻

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samples	excitation peak (nm)	emission peak (nm)	FWHM (nm)	Stokes shift (nm)	decay time (ns)	PLQY (%)
Enrofloxacin	430	544	129	114	2.77	<1
ENR-HCl	443	470	75	27	1.73	2.10
ENR-Pb	365	443	72	78	2.31	9.75
ENR-Bi	422	459	56	37	2.50	9.14
ENR-Sn	435	465	40	30	6.37	19.23





Figure 5. Calculated electronic band structure of ENR-Pb (a), ENR-Bi (b), and ENR-Sn (c) single crystals. Total and orbital-projected partial density of states of ENR-Pb (d), ENR-Bi (e), and ENR-Sn (f) single crystals calculated by DFT-PBE method.

(Figure 3b). There are O-H…Cl, O-H…O, N-H…Cl, C-H…Cl, and C-H…O hydrogen bonds in the structure (Table \$5), three of which have intramolecular hydrogen bonds. Intermolecular hydrogen bonds form a 3D net, which joins the $EnrofloH_2^{2+}$ cations with $BiCl_6^{3-}$ metal-chloride anions, H_2O molecules and Cl⁻ ions. The EnrofloH₂-Sn single crystal also crystallizes in centrosymmetric monoclinic space group $P2_1/c$. The asymmetric part contains one ion of C₁₉H₂₄FN₃O₃²⁺ (Enroflo H_2^{2+}), one Sn⁴⁺ ion, six Cl⁻ ions, and one H_2O molecule (Figure 3c'). The main bond lengths are shown in Tables S6. Each Sn⁴⁺ ion is coordinated by six Cl⁻ ions and forms SnCl₆²⁻ metal-chloride anions, which is isolated from each other (Figure 3c). There are O-H…Cl, O-H…O, N-H…Cl, C-H…Cl, and C-H…O hydrogen bonds in the structure (Table S7), three of which have intramolecular hydrogen bonds. Intermolecular H-bonds form 3D net, which joins the $EnrofloH_2^{2+}$ cations with $SnCl_6^{2-}$ metal-chloride anions and H₂O molecules.

Generally, the emissive organic cations exhibit fluorescence emission arising from singlet state with short lifetime or roomtemperature phosphorescence (RTP) arising from triplet state with long lifetime.^{28–32} The fluorescence spectra of Enrofloxacin powders are shown in Figure S1. Under 430 nm excitation, it shows the broadband yellow emission peaking at 544 nm with the PLQY of <1 %. The fluorescence lifetime of the emission is determined by time-resolved methods (Figure S2). The average lifetime of the emission at room temperature is 2.77 ns based on double exponential-fitting. The room temperature optical properties of as-synthesized metal halides are investigated in detail, as shown in Figure 4. The photoluminescence excitation (PLE) spectra of all three compounds show broad excitation bands from 300-450 nm. Under the excitation at 365 and 422 nm, ENR-Pb and ENR-Bi single crystals show the blue emission peaking at 443 and 459 nm with the Stokes shift of 78 and 37 nm and a FWHM of 72 and 56 nm, respectively (Figure 4a). Under the excitation at 435 nm, the ENR-Sn single crystal shows the blue emissions peaking at 465 nm (Figure 4b,c) with the Stokes shift of 30 nm and a FWHM of 40 nm (Figure 4e,f). The lifetime of ENR-Sn at room temperature is 6.37 ns based on single exponentialfitting. Especially, the average lifetime of ENR-Pb and ENR-Bi at room temperature are 2.31 and 2.5 ns based on double exponential-fitting, respectively, which may be caused by effective nonradiative energy transfer. It is consistent with the observed low PLOYs of ENR-Pb and ENR-Bi single crystals. We attribute these blue emissions to the singlet state fluorescence emission of EnrofloH2²⁺ cations, because the Enrofloxacin in solution also shows the similar blue emissions.^{20,33} As mentioned above, Enrofloxacin exists in the form of LH2²⁺ (divalent cation) in the strong acidic conditions, in which the maximum emission wavelength is located at 450 nm. When Enrofloxacin exists in the form of LH⁺ (monovalent cation) under the pH condition of 2.45-4.23, the maximum emission wavelength is still at 450 nm.



Figure 6. Chemical structure of $\text{EnrofloH}_2^{2^+}$ cations in ENR-Pb (a), ENR-Bi (b), and ENR-Sn (c) single crystals. The dotted lines (a)–(c) show the type of hydrogen bonding. Layered stack of $\text{EnrofloH}_2^{2^+}$ in ENR-Pb (a), ENR-Bi (b), and ENR-Sn (c) single crystals. The dotted lines (d)–(f) show the shortest distance of two adjacent benzene rings in $\text{EnrofloH}_2^{2^+}$ cations.

When the pH value is larger than 4.23, Enrofloxacin exists in the form of L (bipolar), and the maximum emission wavelength gradually shifted to 445 nm with the decreased fluorescence intensity. When the pH value is more than 12.28, Enrofloxacin exists in the form of L^{-} (anion), the maximum emission wavelength is still at 445 nm, but the fluorescence intensity decreased and nearly disappeared with the increase of the pH value.³³ One can conclude that the divalent cation species only exist in extremely acidic conditions. Therefore, we analyze the optical properties of Enrofloxacin in HCl solution with the concentration of 11-12 mol/L (ENR-HCl); the fluorescence spectra of ENR-HCl are shown in Figure S3. Under the excitation at 443 nm, ENR-HCl shows the blue emission peaking at 470 nm with the Stokes shift of 27 nm and a FWHM of 75 nm, and the PLQY of 2.11 %. The detailed optical properties of Enrofloxacin, ENR-HCl, and hybrid metal halides are summarized in Table 1. It is of interest to note that the PL properties of EnrofloH2²⁺ organic cations have significant distinctions in solution and in different single crystals, such as the maximum excitation and emission wavelength, FWHM, and PLQY. Especially, the PLQY of ENR-Sn single crystal reaches 19.73 %, and the FWHM of the corresponding emission peak is only 40 nm. We also analyze the effect of excitation wavelength on the fluorescence emission for these three single crystal compounds (Figures S5-S7). The emission wavelengths remain unchanged with the change of excitation wavelengths, indicating that there was only one luminescence center in these single crystal compounds.

To further verify the PL origin of these as-synthesized metal halides, we analyze the electronic band structures and density of states (DOS) of these three single crystal compounds by density functional theory (DFT). As shown in Figure 4, ENR-Pb, ENR-Bi, and ENR-Sn all have the indirect bandgaps with 2.23, 2.61, and 2.26 eV (Figure 5a-c), respectively. The valence band maximum (VBM) of these compounds are all made up of $EnrofloH_2^{2+}$ -p and Cl-6p orbitals, and the

conduction band minimum (CBM) are all composed of Enroflo H_2^{2+} -p (Figure 5d,e), which indicated that the center metal ions have no contribution to the frontier orbital in these hybrid metal halides. In other words, the fluorescence emissions of all synthesized compounds come from the $EnrofloH_2^{2+}$ organic cations, which is consistent with observed optical characteristic. In addition, it also can be found from the synthesis processes of these compounds that all of the single crystal compounds are composed of protonated organic cations and metal-chloride anions. Therefore, the fluorescence emissions of these single crystal compounds belong to the corresponding organic cations with certain degree of protonation. Figure 6 shows the chemical structures of EnrofloH2²⁺ cations in ENR- Pb, ENR-Bi, and ENR-Sn, which are used to analyze the degree of protonation of Enrofloxacin in targeted compounds. The organic components of these single crystal compounds have the same degree of protonation, including 3-position nitrogen (N3) in the piperazinyl group and 1-position carbonyl oxygen (O1) in the quinoline ring (Figure 6a-c), which is similar to other ionic compounds composed of ciprofloxacin and Bi(III) chloride anions.^{34,35} In neutral and acid solutions, according to the different pH conditions, the fluoroquinolones (FQ) might exist in four protonation species, namely, dication FQH_2^{2+} , cation FQH^+ , zwitterion zFQ, and neutral species nFQ_{2}^{36} The FQH₂²⁺ only exist in extremely acidic conditions, which is consistent with the synthesized conditions of these three compounds in acid conditions with the concentration of HCl solution of 11–12 mol/L. In addition, FQH_2^{2+} can be protonated either at the piperazine ring^{37,38} or at carbonyl oxygen (O1) or even through the atom of pyridone ring (N1).³⁹ The results of the DFT calculation showed that the FQH₂²⁺-O1 form is the most stable isomer in the gas phase and in water.³⁶ These results are also consistent with the observed protonation forms of EnrofloH₂²⁺ cations in our synthesized compounds.

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Figure 7. Energy level diagram of ENR-Pb (a), ENR-Bi (b), and ENR-Sn (c) single crystals. The S_n represent the high energy singlet states, the S_1 represent the low energy singlet state, the S_0 represent the ground state, and the IC represent the internal conversion.

Although Enrofloxacin has the same degree and form of protonation in these three compounds, the optical properties of protonated organic cations in these as-synthesized metal halides are significantly discrepant. As shown in Table 1, the maximum emission wavelength of these as-synthesized metal halides is red-shifted from 443 to 459 to 465 nm by changing metal-chloride anions from $Pb_2Cl_6^{2-}$ to $Bi_2Cl_{10}^{4-}$ to $SnCl_6^{2-}$. Moreover, the PL of ENR-Sn has the smallest Stokes shift of 30 nm and the FWHM of the corresponding emission peak is low to 40 nm and the PLOY is up to 19.23 %. One can conclude that under the same protonation form, the other factors should be involved, which greatly affect the fluorescence emissions of EnrofloH22+ in synthesized single crystal compounds. As mentioned above, the PL of emissive organic cation is greatly affected by their local environment. The inter- and intramolecular hydrogen bonds and intermolecular π stacking interactions can form a more restraining effect and improve the rigid degree of organic cations, causing the red shift or blue shift of the spectrum and the variation of FWHM of corresponding emission peak. There are both interand intramolecular hydrogen bonds in three single crystal compounds. The intermolecular hydrogen bonds joint the EnrofloH22+ with metal-chloride anions forming a 3D net, enhancing the rigid degree of EnrofloH22+. Differently, in ENR-Pb, the proton of the carbonyl bound oxygen (O1) prefers to combine with the oxygen in isolated H₂O molecule rather than the carboxyl group bound oxygen (O3), forming an intermolecular hydrogen bond (Figure 6a). Conversely, in ENR-Bi, the proton of the carbonyl bound oxygen (O1) is combined with carboxyl group bound oxygen (O3), forming an intramolecular hydrogen bond (Figure 6b). Moreover, both intra- and inter- molecular hydrogen bonds are observed in ENR-Sn (Figure 6c), which come from the combination of the proton of the carboxyl group bound oxygen (O2) and oxygen in an isolated H₂O molecule. In addition to hydrogen bond interactions, $\pi - \pi$ stacking interactions also play an important role in improving the rigidity of $EnrofloH_2^{2+}$ cations. Generally,

the organic cations in solution have more considerable flexibility. However, when liquid organic cations are transformed into solids forming highly ordered crystals, which have decreased flexibility. Furthermore, the denser packing of EnrofloH₂²⁺ cations in single crystal compounds significantly reduces their flexibility due to $\pi - \pi$ stacking interactions. Figure 6d-f shows the crystal packing of EnrofloH₂²⁺ cations in these single crystal compounds. Enroflo H_2^{2+} cations show the two infinite stacks (Figure 6d), and the offset face-to-face $\pi-\pi$ stacking of phenyl rings are observed in the ENR-Pb compound, here the shortest distance between adjacent phenyl rings is 3.530 (7) Å. The Enroflo H_2^{2+} cations also show the layered stack and the shortest distances between adjacent phenyl rings are 5.032 (6) Å in the ENR-Bi compound (Figure 6e), which are greater than the 4 Å,⁴⁰ so there are no $\pi - \pi$ stacking interactions. Moreover, EnrofloH₂²⁺ cations in ENR-Sn are stacked in layers, and adjacent layers are arranged vertically. The phenyl rings in $EnrofloH_2^{2+}$ cations show the offset face-to-face $\pi - \pi$ stacking (Figure 6f), and the shortest distances between adjacent phenyl rings are 3.735 (2) Å. The detailed parameters of the $\pi-\pi$ interactions in ENR-Pb and ENR-Sn single crystal compounds are listed in Table S8. The energy level diagrams of ENR-Pb, ENR-Bi, and ENR-Sn single crystal compounds are shown in Figure 7. Under the excitation at UV light, the ground state electrons are excited to a high energy singlet state (S_n) , which undergoes the internal conversion (IC) to the lower energy singlet state (S_1) and then returns to ground state (S₀) to generate Stokes-shifted emissions with nanosecond decay times. However, due to the diverse local environment, the optical properties of Enro $floH_2^{2+}$ cations in these three single crystal compounds are different. Thus, because of the coupling effect of hydrogen bonds and $\pi - \pi$ stacking interactions, the emission of EnrofloH₂²⁺ cations in ENR-Sn shows the smallest Stokes shift of 30 nm and the FWHM of the corresponding emission peak is low to 40 nm and the PLQY of up to 19.23 % with the best performance in the studied series.

In summary, we demonstrated an all-purpose strategy for modulating the optical properties of emissive organic cations by assembling them with metal-chloride anions. The single crystal bulk assembly of EnrofloH2²⁺ organic cation and different metal-chloride anions $(Pb_2Cl_6^{2-}, Bi_2Cl_{10}^{4-}, SnCl_6^{2-})$ can draw into additional interactions, improve the optical properties of $EnrofloH_2^{2+}$ organic cation, including modified excitation and emission wavelength, FWHM, and PLQY. The local structures and optical properties studies indicated that the hydrogen bond and $\pi - \pi$ interactions can provide the rigid environment for the organic cation, which decrease the flexibility of the organic cation. Thus, the fluorescence emissions of the EnrofloH2²⁺ organic cation in hybrid metal halides have wide differences compared with pure organic cations in solutions. Our work not only demonstrates the role of metal-chloride anions in photoluminescence modulation of emissive organic cations in hybrid metal halides materials for future emerging optoelectronic applications but also provides a fundamental way for designing multifarious optical functional materials.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpclett.1c00182.

Experimental section, Figures S1–S7 showing excitation and emission spectra, decay curves, and Tables S1–S8 providing main parameters of single crystal structure and the π - π interactions in ENR-Pb, ENR-Bi, and ENR-Sn single crystals (PDF)

Crystallographic information of $(EnrofloH_2)Pb_2Cl_6 H_2O$ (CIF)

Crystallographic information of $(EnrofloH_2)BiCl_5 \cdot Cl \cdot 2(H_2O) \cdot H_3O$ (CIF)

Crystallographic information of $(EnrofloH_2)SnCl_6 H_2O$ (CIF)

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Notes

The authors declare no competing financial interest.

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