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White light emission in low-dimensional perovskites

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Low-dimensional perovskites are rapidly emerging for their distinctive emission properties, consisting in ultrabroad and highly Stokes shifted luminescence with pure white light chromaticity, which makes them very attractive for solution-processed light-emitting devices and scintillators. To foster the design of new materials and their device applications, it is timely to review the relation between perovskite structural properties and photophysical phenomena underlying their unique light emission characteristics. From a number of recent works, it has emerged that broadband emission properties in metal halide frameworks are very common, stemming from the self-localization of small polaron species at specific sites of the inorganic lattice, with a wide energy distribution. This review aims to provide an account of the current understanding of the photophysical processes underpinning luminescence broadening and highly efficient emission in various classes of low-dimensional metal-halide frameworks, and to highlight their potential for solution-processed optoelectronic device applications. The discussion will additionally establish a wider perspective on the role of intrinsic and extrinsic self-trapping, polarons formation and their effect on charge generation and transport in low-dimensional perovskites.

1 Introduction

Three-dimensional (3D) hybrid perovskites have rapidly established new paradigms for solution processed photovoltaics (reaching power conversion efficiencies up to 22%) and light emitting devices.1–3 At an equally fast pace, low-dimensional perovskites are opening new frontiers for solution-processable optoelectronic devices.4,5 Primarily based on self-assembled metal-organic frameworks with highly tunable composition and reduced dimensionality (2D, 1D or 0D), this class of materials adds synthetic versatility to the standard 3D perovskite structure, providing distinctive advantages for light-emitting applications achievable by structural design:6 strong exciton localization effects result in unusually large Stokes shift,7 which reduces luminescence self-absorption and thermal quenching; ultrabroad luminescence spectral width yields purely white chromaticity from a single emissive material;8 the hydrophobicity of the templating organic cations significantly improves material stability in ambient conditions;9 furthermore, the presence of heavy ions provides large absorption cross section for ionizing radiation.10 These features make low-dimensional perovskites particularly appealing for white light emitting devices (diodes, transistors and lasers) and large area X-ray scintillators for solid-state lighting, displays, medical, and security applications. Recent progresses in the synthesis of new classes of low-dimensional perovskites are revealing widespread broadband emissive characteristics. In these systems, the ability to govern optoelectronic properties through rational synthetic design is conditional to the deep understanding of their unusual photophysical properties. This review aims at consolidating the recent understanding of the processes underpinning luminescence broadening and highly efficient emission in a variety of low-dimensional hybrid perovskites, with foreseeable device applications. Specifically, experimental and theoretical studies highlighted the role of charge self-trapping on the emission characteristics of low-dimensional perovskites, impinging the ultrafast formation of emissive color centers at specific inorganic lattice sites. From a broader fundamental standpoint, a unified picture of (small) polaron formation in relation to structural and compositional properties provides a common ground to discuss implications on charge generation and transport in metal halide frameworks with different dimensionalities, beyond their luminescence characteristics.

1.1 From 3D to low dimensional perovskites. Three dimensional (3D) perovskites have the general formula AMX₃, where X = F⁻, Cl⁻, Br⁻, I⁻, and A is a divalent metal or organic cation.¹¹ This structure is characterized by a continuous 3D network of corner-sharing MX₆ octahedra, while A is 12-fold coordinated in the cuboctahedral cavities, which limit the size of the cation to be enclosed.¹² The Goldschmidt tolerance factor t = (Rₐ + Rₓ) / (2(Rₐ + Rₓ)) and the octahedral factor µ = Rₘ / Rₓ (Rₐ, Rₓ and Rₘ are the effective ionic radii of the corresponding ions) help to predict whether a’ would fit in the 3D perovskite structure, that is stable when 0.8 < t < 1 and 0.442 < µ < 0.895.¹³,¹⁴ Big cations disrupt the 3D network and form low dimensional structures where BX₆ octahedra only partially share their corners with neighbors. Low dimensional perovskites can be derived by slicing the parental 3D perovskite along different orientations (Fig. 1), which significantly

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perovskites are formed, characterized by chains of MXn units sharing two corners in equatorial position (Fig. 1b, left). For m ≥ 2, the resulting 2D perovskites are characterized by rippled inorganic layers terminated by two undercoordinated halides from each MX6 octahedron, which therefore ends the layer exposing one edge (Fig. 1b, right). Lastly, <111>-oriented perovskites with formula A′2A3n-1MnX3n+1b form zero-dimensional (0D) structures (when q = 1) made of isolated MX6 octahedra (Fig. 1c, left). The 2D perovskite (q = 2; Fig. 1c, right) is intrinsically different from the previous two classes, since B must be a trivalent cation to maintain electroneutrality. Here, the inorganic sheets are made of octahedra sharing three corners and terminating the inorganic slabs exposing one face, therefore contributing 3 undercoordinated halides to the terminating surface.\(^a\) Thanks to the enormous synthetic versatility, a large number of alternative metal halide frameworks can be derived from the archetype structures shown in Fig. 1, for instance forming combinations of <100> and <110> perovskite frameworks or inducing different connectivity of corner-, edge- and face-sharing octahedra.\(^b\)

1.2 Distortions of the perovskite lattice. The band structure, emissive properties and defect formation energy of perovskites are significantly affected by the connectivity of the inorganic motif.\(^c\) The perovskite structural properties can be evaluated using some relevant parameters:

\[ M\text{-X-M angle and M-X bond length. While M-X length has generally a secondary effect on the absorption properties of the perovskite, the} \]

\[ \text{M-X-M angle has a major impact on the optical properties.} \]

Considering the case of <100>-oriented 2D perovskites, deviation of the M-X-M angle from the ideal geometry (180°) implicates distortions of the inorganic frameworks involving in-plane (\(\theta_{\text{in}}\)) and out-of-plane (\(\theta_{\text{out}}\)) tilt of the MX6 octahedra.\(^d\) The greater the tilt, the poorer the overlap between the metal and halide orbitals, widening the band-gap and causing the blue-shift of the absorption onset.\(^e\)

**Octahedral distortion.** Distortions of the MX6 octahedra combine deformations from the ideal Oh symmetry of M-X bond lengths (\(d_{\text{1}}\)), X-M-X angles (\(\alpha_{\text{i}}\)), edges (X-X distances) and volume (\(V\)), which can also affect the luminescence properties of the material. For example, in the series \((C_{x}H_{y}NH_{z})_{2}PbBr_{6}\), \((C_{x}H_{y}CH_{z}NH_{w})_{2}PbBr_{6}\) and \((C_{x}H_{y}C_{z}H_{w}NH_{z})_{2}PbBr_{6}\), the PL efficiency and lifetime decrease with the distortion of PbBr\(_6\) octahedra as a consequence of the increase of the reduced mass of the excitons in relation to the structural distortion.\(^f\) Octahedral distortion can be quantitatively evaluated by combining the octahedral angle variance \(\sigma_{\alpha_{\text{i}}}^2\) and octahedral elongation \(\lambda_{\text{oct}}\), which are defined as:

\[ \sigma_{\alpha_{\text{i}}}^2 = \frac{1}{12} \sum_{i=1}^{12} (\alpha_{\text{i}} - 90)^2 \]  

\[ \lambda_{\text{oct}} = \frac{1}{6} \sum_{i=1}^{12} (d_{i/d_{0}})^2 \]
where $d_0$ indicates the center-to-vertex distance of a regular polyhedron of the same volume. Alternatively, the octahedral distortion is also expressed by the distortion parameter $\Delta_d$:

$$\Delta_d = \frac{\sum_{i=1}^{6} (d_i - d_{av})^2}{d_{av}^2}$$  \[3\]

where $d_{av}$ is the average M-X bond length.

Octahedral connectivity. The connectivity of the MX$_6$ units is affected by the perovskite orientation, dimensionality and by the number of shared halide atoms between adjacent octahedra. The connectivity gradually increases from corner to edge and face-sharing octahedra, which results in band-gap widening and can modulate the confinement effects in the low dimensional structures.\textsuperscript{19,25,38}

2 Broadband emission in low dimensional perovskites

Low-dimensional perovskites behave like confined nanostructures,\textsuperscript{6,15} where electronic and dielectric confinement arises from differences in band-gap and polarizability of the inorganic and inorganic components. These conditions increase significantly the electron-hole Coulomb interaction (through image charge effect) and the exciton binding energy (EBE) up to few hundreds of meV,\textsuperscript{39,40} so that exciton population may become predominant compared to free-charges even at low excitation densities.\textsuperscript{41} This is in net contrast to 3D perovskites, where high dielectric constant and efficient Coulomb screening confers a remarkably low EBE (<50 meV).\textsuperscript{42,43} The resulting excitonic properties should then drive the characteristic photophysics, with the typical sharp excitonic absorption and extremely narrowband photoluminescence (full-width at half maximum FWHM ~20 nm) characteristic of free-excitonic states.\textsuperscript{40,44,45} However, many low dimensional perovskites show...
opposite behavior,\textsuperscript{46,47} with ultra-broadband (FWHM beyond 100 nm) and highly Stokes-shifted photoluminescence (>1 eV) arising from intra-gap states with wide energy distribution.\textsuperscript{7,48} Perovskite dimensionality, connectivity and structural distortions contribute to shape the energetic landscape in these materials, ultimately determining their emissive properties; the following paragraphs summarize a wide range of experimental evidences to rationalize such structure/photophysics relationship.

2.1 2D perovskites. <100>-oriented 2D perovskites display a complex trend in their photoluminescence since the balance between narrowband and broadband room-temperature emission appears to be strongly related to their structural properties. While structures with close-to-ideal geometries show exclusively free-excitonic emission (e.g., BA\textsubscript{4}PbBr\textsubscript{4} where BA = n-butylammonium), more distorted structures have a prominent contribution from broadband PL (e.g. (EDBE)\textsubscript{2}PbCl\textsubscript{4} (Fig. 2a), while EDBE = 2,2-(ethylenedioxy)bis(ethylammonium)).\textsuperscript{7,48}\textsuperscript{7} While this has been mainly studied in lead-based perovskites, similar behaviour has also been observed in the cadmium-based (\textit{C}_{6}H_{12}N\textsubscript{3})\textsubscript{2}CdBr\textsubscript{4}.\textsuperscript{49} In intermediate cases, the combination of narrow/broadband PL with comparable intensity can be observed. This is the case of (cis-CyBMA)\textsubscript{2}PbBr\textsubscript{3} (cis-CyBMA = 1,3-bis(methylamino)cyclohexane) (Fig. 2b,c) and (2meptH\textsubscript{2})\textsubscript{2}PbBr\textsubscript{4} (2mept = 2-methyl-1,5-diaminopentane), which have PbBr\textsubscript{4} octahedra with close-to-ideal geometry but features a relevant octahedral tilt with Pb-Br-Pb average bond angle <150° (Fig. 2b).\textsuperscript{32,50,52} Recently, M. D. Smith \textit{et al.} rationalized the structure/optical properties relationship by examining the ratio between the intensities of the broad emission (I\textsubscript{b}) and the narrow emission (I\textsubscript{n}) in a set of eight <100>-oriented bromide perovskites.\textsuperscript{45,53} Within this dataset, it was found that the relative intensity of the broad emission (given by the ratio I\textsubscript{b}/I\textsubscript{n}) increases with the increase of out-of-plane distortion of the Pb–(μ-Br)–Pb angle, with the exception of (CEA)\textsubscript{2}PbBr\textsubscript{4} (CEA = cyclohexylammonium) which shows room temperature broadband emission despite the absence of out-of-plane tilt. On the other hand, a univocal correlation of the I\textsubscript{b}/I\textsubscript{n} ratio with the octahedral distortion was not clearly identified.\textsuperscript{27,53} The structural properties play a similar role also in the multidimensional series EA\textsubscript{2}Pb\textsubscript{10}−xCl\textsubscript{x} (Fig. 2d); here distortions of the octahedral coordination as well as octahedral tilt increase from x = 0 to x = 10, allowing to reach white PL in EA\textsubscript{2}Pb\textsubscript{10} (Fig. 2d) compared to the blue emission of EA\textsubscript{2}Pb\textsubscript{8}I\textsubscript{10}.\textsuperscript{13}

The corrugated <110>-oriented perovskites usually endures important steric hindrance effects from the templating cations which result in highly distorted geometries. Moreover, the presence of two undercoordinated halides at the organic/inorganic interface large lattic rearrangements easing the formation of photoinduced defects or structural rearrangements. As a consequence, all the reported bromide-based <110> perovskites show a prominent contribution of the broadband emission at room temperature. These include (API)\textsubscript{2}PbBr\textsubscript{4} and (AEU)\textsubscript{2}PbBr\textsubscript{4} (API = N-(3-aminopropyl)imidazole and AETU = -(aminoethyl)lisothiourea),\textsuperscript{21,54} N-MEDA)PbBr\textsubscript{4} [N-MEDA = N1-methylthene-1,2-diammonium].\textsuperscript{55} and (EDBE)\textsubscript{2}PbBr\textsubscript{4}.\textsuperscript{8} The latter shows photoluminescence quantum yield of 9%, which is the highest reported among bulk 2D perovskites. Similar broadband emission was reported by L. Mao \textit{et al.} in α-(DMEN)Pb\textsubscript{2}Br\textsubscript{6} [DMEN = 2-(dimethylamino)ethylamine], featuring a 3×3 corrugated structure of the <110>-type (Fig. 2e). In comparison with other 2D bromide perovskites based on DMA\textsubscript{2}Pb\textsubscript{4} (DM = 3-(dimethylamino)-1-propylamine and DMABA = 4-dimethylaminobutylamine, the authors identified a correlation between increased octahedral distortion and PL broadening (Fig. 2f).\textsuperscript{34} Despite some variability between different compounds, the broadband emission in <100> and <110>-oriented bromide-based 2D perovskites shares some common characteristics: it is typically centered in the range 500-560 nm, largely Stokes-shifted (>1 eV), FWHM up to 160 nm and photoluminescence decay varying across the emission spectrum (Fig. 2g) reported in the range 0.4-15 ns, pointing to the presence of multiple emissive color centers.\textsuperscript{7} On the other hand, iodide-based perovskites show a slightly different scenario. For example, the <110>-oriented iodide perovskite (EDBE)\textsubscript{2}PbI\textsubscript{4} does not show white light emission at room temperature, however it is also characterized by a considerable broadening of the excitonic emission (FWHM=70 nm) which correlates with the increased structural distortion compared to purely free-excitonic emitting iodide-based 2D perovskites (Fig. 2h).\textsuperscript{35} At low temperature, largely Stokes shifted broadband luminescence centered around 600-625 nm was also observed in alkylammonium-based iodide perovskites such as (NBT)\textsubscript{2}Pbl\textsubscript{4} (NBT = n-butylammonium; Fig. 3i), DA\textsubscript{2}Pbl\textsubscript{4} (DA = dodecylammonium) and HA\textsubscript{2}Pbl\textsubscript{4} (HA = hexylammonium). Here, the broadband emission was measured only below 200 K and no evident correlation between octahedral tilt and appearance of broadband emission was observed.\textsuperscript{36,56}

In <111>-oriented perovskites, the presence of three undercoordinated halides at the organic/inorganic interface further ease structural distortions and coordination rearrangements, suggesting that should also be prone to broadband emission. Being based on trivalent cations Sb\textsuperscript{3+} and Bi\textsuperscript{3+},\textsuperscript{57,58} they also present some differences compared to the previous two perovskite types (e.g. the PL is usually more red-shifted), although only few examples are available. While Rb\textsubscript{2}Bi\textsubscript{3}S\textsubscript{3} shows no PL response (probably due to predominance of non-radiative recombination pathways), Cs\textsubscript{2}Sb\textsubscript{3} films display highly Stokes shifted PL peaked at 646 nm, FWHM of ~60 nm and the complete lack of free-exciton emission.\textsuperscript{59}

Even though structural deformations appear to strongly impact on the luminescence characteristics of this class of perovskites, a unified model describing how the structural properties shape the energetic landscape of 2D perovskites with different orientation and composition is still missing. Similarly, the reports supporting the relative contributions of octahedral distortion or tilt to the broadening process are sometimes conflicting, leaving space for further investigations. Synthesis of new materials and photophysical characterization should provide new insights to guide the crystal engineering of layered perovskites with controlled optical properties.
2.2 1D and 0D perovskites. Z. Yuan et al. discovered that the cation N, N'-dimethylethylenediammonium (DMEDA) templates the growth of the pseudo-1D perovskite (DMEDA)PbBr₄ in which lead bromide wires are composed of double edge-sharing PbBr₆ octahedra, surrounded by a shell of DMEDA²⁻ cations (Fig. 3a). The material could be synthetized in form of bulk crystalline needles having similar emission to that characteristic of layered perovskites: highly Stokes-shifted (1 eV), cold bluish white light centered at 475 nm with PLQY up to 18-20% and lifetime of 37.3 ns at room temperature (Fig. 3a). The corresponding chloride perovskite (DMEDA)PbCl₄ displayed excitation-dependent broad photoluminescence, attributed to the competition between different emission centres. The use of DMEDA with Sn²⁺ was also reported to form the 0D perovskites (DMEDA)₄SnBr₆ and (DMEDA)₃SnI₆ with orange and red emission, FWHM >100 nm and Stokes shift ~200 nm. In particular, the iodide-based perovskite showed an exceptionally long PL lifetime of 1.1 µs and extremely high PLQY up to 95% at room temperature, which was ascribed to the increased quantum confinement in the 0D structure. PL enhancement by means of decreased dimensionality have also been exploited to obtain warm white-light with remarkably high PLQY (up to 86%) in lead-free double perovskites such as Cs₂AgInCl₆. Despite being formed by a continuous metal halide network, the presence of alternating AgCl₆ and InCl₆ octahedra brakes the continuity of orbital overlap resulting in a reduced 0D-like electronic dimensionality. Charge confinement within single octahedral units was further enhanced by means of Na⁺ and Bi³⁺ doping. Hexamethyldiammonium (HMTA) was used for the synthesis of (HMTA)₂PbBr₄ which forms a bulk assembly of 1D lead bromide nanotubes: six dimers of face-sharing lead bromide octahedra (Pb₅Br₁₂) connect at the corners forming a ring with inner radius of 4Å which extends in one dimension to form the nanotube (Fig. 3b). Here, lead bromide octahedra are highly distorted, with Pb-Br bonds in the range 2.873-3.163 Å and Br-Pb-Br angle between 77.25° to 113.15°. Photoluminescence between 250-400 nm resulted in broadband emission (FWHM ~ 158 nm) peaked at 580 nm with PLQY ~ 7% and long lifetime of 106 ns. Cooling at 77K allowed to resolve three distinct emission peaks centered at 420, 480 and 625 nm with lifetime of 7, 4.0 and 1.8 µs, respectively, suggesting that multiple species are involved in the emission process (Fig. 3c,d). Ethylene diammonium and piperazine were also shown to template pseudo 1D and 0D structures with broadband emission.

2.3 Exotic metal halide frameworks. Photoluminescence broadening was also shown in various examples of more complex lead halide frameworks, suggesting a relation with the nature of the metal halide bonds rather than to intrinsic properties of the perovskite structure. For example, trimethylphenylammonium (TMPA) and a series of sulfonium cations including trimethylsulfonium (TMS) and butane-1,4-bis(dimethylsulfonium) (1,4-BDDMS) were shown to template structures in which isolated trimers or pentamers of face-sharing lead bromide octahedra are surrounded by the organic cations. These materials have similar optical properties, exemplified by the case of (1,4-bddms)₃PbBr₁₂ in Fig. 4a,b: excitonic properties comparable to those of standard 2D perovskites, highly Stokes-shifted (> 1.7 eV) red photoluminescence centered at 690 nm and FWHM ~ 0.7 eV. Such broadband red luminescence appears to be characteristic of this type of connectivity of the inorganic framework, where octahedra are connected through one of their faces. Interestingly, white light luminescence was also achieved in the 3D haloplumbate frameworks (H₂DABCO)[PbCl₄] and (H₂O)[Et₂DABCO]₆[Pb₂Cl₆]₄, where DABCO = 1,4-diazabicyclo[2.2.2]-octane and Et = ethyl (Fig. 4c,d). In this case, white light emission was obtained by combining blue photoluminescence of the organic cation with broad yellow emission originating from the lead chloride network, respectively centered at 455 and 585 nm in (H₂DABCO)[PbCl₄] and (H₂DABCO)[Pb₂Cl₆]₄. Despite the modest PLQY (1-2.5%), a very high color rendering index (CRI) value was achieved (96) together with electric conductivity up to 2.83 x 10⁻⁴ S cm⁻¹.

S. Z. Zhuang et al. recently synthesized a series of rare cationic layered lead halides with formula [PbₓX₄]ₙ⁺[CO₃(CH₂)₂CO₃]⁻ (X = F, Cl, and Br) by hydrothermal reaction, where corrugated cationic lead halide sheets are held in place by elongated adipate anions (Fig. 4e,f). These materials differ from standard 2D perovskites since the carboxylate is directly coordinated to the Pb²⁺ centres through covalent bonds. In the fluoride compound the Pb²⁺ is further coordinated with four fluorine atoms forming a distorted square-planar geometry, while the chloride and bromide compounds form PbX₄ units which are analogous to half of the PbX₄ octahedra of typical 2D layered perovskites. Set aside the contribution of the uncoordinated axial ligands, the PbX₄ units of the Cl and Br-substituted materials showed remarkable octahedral distortion, with...
3 Origin of broadband emission

The observation of such a dominant contribution of broadband, highly Stokes shifted emission in low-dimensional perovskites have stimulated the scientific community to identify the physical processes underlying this phenomenon. Mechanistic studies performed on several metal halides reveal common characteristics which point to the formation of self-trapped excitons, which define the emissive intra gap states (see paragraph 3.1 for a detailed description).71 In this regard, the main observations include: i) Emission bandwidth and PLQY are unrelated to the perovskite crystal size remaining substantially unchanged in films, powders and nanocrystals, excluding the involvement of surface/edge states in the emission process.8,70,72 ii) The integrated PL intensity of broadband emission linearly increases with the excitation intensity over several orders of magnitude without showing saturation, ruling out the contribution of permanent defects (particularly in Cl and Br perovskites)8,62,63,67 iii) The PL temperature dependence is consistent with the presence of an energy barrier for the self-trapping process, which is expected given the structural rearrangement necessary to induce charge localization.73

The contribution of broadband PL increases at lower temperature respect to free-excitonic emission, accompanied by an increase of the overall luminescence efficiency.31,50 This result from the lower probability of exciton de-trapping jointly with the reduction of the impact of non-radiative recombination.

When temperature is further reduced (usually below ~80 K), the free excitons become progressively unable to overcome the energy barrier to undergo self-trapping, and the relative contribution of free-excitonic emission starts to increase again.47,52,53,73 iv) Transient absorption measurements revealed the formation of an unstructured photoinduced absorption band covering the entire visible spectral region, pointing to the formation of multiple emissive color centers distributed throughout the perovskite band-gap. On the contrary, narrowband emitters show bleaching features below the excitonic region, pointing to the formation of multiple emissive color centers.

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characteristic luminescence and absorption line-shapes. Where excitonic polarons must be considered to account for their lattice deformations have been identified in several 2D perovskites, highly polar lead halide framework. Recently, strong photoinduced confinement sustain tightly bound excitons which interact with the density distributions for (EDBE)PbCl$_4$ and conduction band minimum (CBM), and corresponding charge density distributions within the central [PbBr$_6$]$^{2-}$ octahedron for positive and negative small polaron (Pb$^{2+}$/Pb$^{3+}$) states; reproduced from ref. 85 Copyright 2017 American Chemical Society. d) Transient absorption spectra of Cs$_2$PbBr$_6$ with different delay times using 310 nm excitation and charge density distributions within the central [PbBr$_6$]$^{2-}$ octahedron for positive and negative small polaron in Cs$_2$PbBr$_6$ supercell; reprinted with permission from ref. 86 Copyright 2017 American Association for the Advancement of Science.

In low-dimensional systems, the coulombic and dielectric confinement sustain tightly bound excitons which interact with the highly polar lead halide framework. Recently, strong photoinduced lattice deformations have been identified in several 2D perovskites, where excitonic polarons must be considered to account for their characteristic luminescence and absorption line-shapes. The strong exciton-phonon coupling can ease exciton self-trapping, leading to the photoinduced generation of small polarons in which the excess charges are spatially confined to one crystal unit cell or less. In this case, the short-range deformation potential due to the changes in local bonding related to the excess charge, plays the dominant role. Due to their strongly confined nature, the formation of such polaronic species is highly dependent on the local structural properties of the crystal lattice, similar to the case of halide-bridged mixed-valence transition metal linear chain complexes. These small polarons can be viewed as self-trapped holes or electrons, as often observed in alkali, alkaline-earth, and perovskite-structure halides, and demonstrated in early work on lead halides (PbBr$_2$ and PbCl$_2$) using electron paramagnetic resonance (EPR). Because of the strong lattice deformation associated with the trapping process, the excited state is shifted compared to the ground state, and radiative decay results in broad and highly Stokes shifted luminescence. In a similar way, in 2D hybrid perovskites the broadband emission can be interpreted as the superimposed, homogeneously broadened luminescence from a series of multiple polaron-exciton states. As shown in Fig. 5a, we have simulated the photogeneration of small polarons and charge self-trapping effects in 2D hybrid perovskites by exerting local perturbations onto the 2D perovskite periodic models. By shortening selected bond lengths (i.e. Pb-Pb, Pb-X and X-X) to create the local structural deformations, we found that i) the Pb-Pb dimerization leads to the formation of a self-trapped electron at Pb$^{2+}$ sites; ii) the hole density is highly localized at the Cl-Cl or Br-Br pairs, showing the formation of V$_k$ centers (Cl$_2$ and Br$_2$); and iii) holes localize at a single Pb atom, leading to the formation of Pb$^{3+}$ centers coupled with the lattice deformation. Therefore, the strong interaction between charge carriers and phonons can yield the formation of self-trapped electron (STEL, Pb$^{3+}$) and self-trapped hole (STH, Pb$^{2+}$ and X$_2$) states. These STEL and STH, consisting of charges trapped at specific sites of the inorganic lattice by their own distortion field, define a series of intra-band polaronic emissive states which decay radiatively with broadband emission (see Fig. 5b). We have also considered a molecular-cluster model to characterize the actual small polaron distributions in the 2D hybrid perovskites. The resulting charge density distribution confirms that the local structural change induces the formation of a localized, positive small polaron (Pb$^{3+}$ self-trapped hole). The strongly localized additional electron density surrounding Pb$^{2+}$ induced large repulsive interactions between adjacent Pb and Cl atoms, thus markedly deforming the local geometry of the 2D perovskite clusters. Similarly, the formation of V$_k$ centers and electron polarons was proposed to explain the scintillation properties of Cs$_2$HfCl$_6$. Recently, X. Wang et al. showed by means of density functional theory calculations that reduced electronic dimensionality plays a crucial role in the formation of self-trapped excitons. Although several species can form, it was proposed that self-trapped excitons connected to Jahn–Teller-like octahedral distortion give the main contribution to the broadband luminescence, while all other types of self-trapped excitons act as non-radiative recombination channels. The existence of such two competing mechanisms suggests that Jahn–Teller-like self-trapped excitons must be promoted while suppressing non-Jahn–Teller-like states in order to obtain high photoluminescence quantum yields. This represents a step forward...
the establishment of a general structure-property relationship in systems with different structural dimensionality and accounts for the experimental observation of broadband emission primarily in the most distorted structures.

Broadband emission can also be achieved at the high-energy spectral region in 0D CsPbBr\(_3\) nanocrystals [see Fig. 5c]. However, the origin of such emission is different from 2D hybrid perovskites: the UV emission around 350 nm can be assigned to the allowed optical transition of \(\text{Pb}\text{Br}_6^{2-}\) ions and the low-energy UV emission at approximately 400 nm to the charge transfer state involved in the 0D host lattice (i.e., D-state). In this case, we can effectively tune the Pb\(^{2+}\)ion emissions in the visible spectral region by adopting different 0D perovskite host lattices. The 0D perovskites are also ideal platforms for studying photogenerated small polarons since the complete isolation of the octahedra is expected to strengthen electron-phonon interactions. As shown in Fig. 5d, the femtosecond transient absorption of Cs/PbBr\(_3\) thin films shows a new positive broadband appearing above 530 nm, due to polaron absorption. The hole charge density is highly localized at the central site once the Pb-Br bonds of the central [PbBr\(_6\)]\(^4-\) unit are shortened, leading to the formation of Pb\(^{3+}\) centers coupled with the lattice deformation. The individual [PbBr\(_6\)]\(^4-\) octahedra in the 0D crystal can be easily perturbed by photoexcitation processes, generating polaronic states through structural deformation potentials. These results suggest that the increased electron-phonon coupling in low-dimensional perovskites can facilitate small polaron generation providing important optical fingerprints, like broadband emission and polaron band absorption.

### 3.2 The role of extrinsic defects

The previous paragraph discussed cases of intrinsic self-trapping, purely polaronic effects in a perfect crystal where a carrier is bound within a potential well arising from the displacement of surrounding atoms to new equilibrium positions. However, the presence of permanent defects (e.g., vacancies, interstitial atoms or impurities) can aid the trapping process, and a charge might localize on one of several symmetry equivalent ions neighboring a defect. In some cases, this trapping might retain a polaronic character leading to a phenomenon identified as extrinsic self-trapping. This is the case, for example, of electron trapping at oxygen vacancies in HFO\(_2\), charge trapping in amorphous SiO\(_2\), hole trapping at Mg\(^{2+}\) vacancies in MgO or small polarons in doped LaMnO\(_3\). Given the relevance of defects in determining the photophysical behavior of 3D perovskites, it is likely that extrinsic trapping might play a role also in low-dimensional perovskites.

While Cl and Br based low-dimensional perovskites have been mainly discussed in terms of intrinsic self-trapping, iodide-based 2D perovskites have offered hints for discussion of the role of permanent defects. For example, in (EDBE)Pbl\(_4\), the PL power dependence shows a monotonic reduction, indicating its relation with permanent defects which are saturated at higher excitation densities. Here, ab-initio calculations showed that the partial oxidation of two iodine atoms in proximity of a Pb vacancy can form stable \(V_\Gamma\) centers, consisting in \(I_3\) trimers stabilized by a vicinal lead vacancy (Fig. 6); overall, the process involves the trapping of two holes in proximity of the point defect, further inducing a local structural rearrangement coupled to the charge localization. On the other hand, iodide interstitials have been proposed to play a role in the radiative recombination process leading to the broadband emission in HA\(_3\)Pbl\(_4\) and DA\(_3\)Pbl\(_4\). We note that in <111> 2D and 0D perovskites such as Cs\(_2\)SbI\(_4\) and Cs\(_2\)BiI\(_6\), PL power dependence with both linear and sublinear behavior have been reported. This leaves open questions on the possibility of intrinsic self-trapping, while inviting the reconsideration of the role of extrinsic defects and trap assisted emission in these systems.
Table 1 Reported room temperature luminescence properties of broadband emitting metal halide frameworks. *values calculated from the reported spectra; “av” indicates average lifetimes.

<table>
<thead>
<tr>
<th>Templating Cation</th>
<th>Perovskite</th>
<th>CIE (x,y)</th>
<th>CRI</th>
<th>CCT</th>
<th>Broadband PL lifetime (ns)</th>
<th>PLQY %</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>3D haloplumbate framework</td>
<td>(H2DABCO)(PbCl2)</td>
<td>(0.33, 0.34)</td>
<td>96</td>
<td>5393</td>
<td>18 (λem = 585 nm)</td>
<td>2.5</td>
<td>68</td>
</tr>
<tr>
<td></td>
<td>(H2O)(Et2-DABCO)(PbCl2)</td>
<td>(0.38, 0.31)</td>
<td>88</td>
<td>3496</td>
<td>54 (λem = 690 nm)</td>
<td>1.0</td>
<td>68</td>
</tr>
<tr>
<td>Cationic 3D lead halide</td>
<td>[Pb2Br6][O2C(CH2)3CO2]-</td>
<td>(0.33, 0.48)</td>
<td>56*</td>
<td>5727</td>
<td>1.7 (λem = 530 nm; av)</td>
<td>5.6</td>
<td>70</td>
</tr>
<tr>
<td></td>
<td>[Pb3Br8][O2C(CH2)3CO2]-</td>
<td>(0.25, 0.32)</td>
<td>72*</td>
<td>11967</td>
<td>1.6 (λem = 480 nm; av)</td>
<td>1.8</td>
<td>70</td>
</tr>
<tr>
<td>multidimensional perovskite</td>
<td>(EA)2PbBr4Cl2</td>
<td>(0.30, 0.35)</td>
<td>83</td>
<td>7132</td>
<td>1.21 (av)</td>
<td>-</td>
<td>33</td>
</tr>
<tr>
<td></td>
<td>(EDBE)2PbBr4</td>
<td>(0.33, 0.39)</td>
<td>81</td>
<td>3990</td>
<td>0.04 – 3.8</td>
<td>2.0</td>
<td>5,7</td>
</tr>
<tr>
<td></td>
<td>(AEP)2PbBr4</td>
<td>(0.29, 0.34)</td>
<td>87</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>53</td>
</tr>
<tr>
<td>&lt;100&gt; 2D perovskite</td>
<td>(AEP)2PbBr4</td>
<td>(0.48, 0.45)</td>
<td>28</td>
<td>154</td>
<td>-</td>
<td>-</td>
<td>52</td>
</tr>
<tr>
<td></td>
<td>(CEA)2PbBr4</td>
<td>(0.31, 0.34)</td>
<td>94</td>
<td>6785</td>
<td>1.2 – 2.2</td>
<td>1.6</td>
<td>32</td>
</tr>
<tr>
<td></td>
<td>(PEA)2PbCl4</td>
<td>(0.37, 0.42)</td>
<td>84</td>
<td>4426</td>
<td>3 (λem = 545 nm; av)</td>
<td>&lt;1</td>
<td>72</td>
</tr>
<tr>
<td></td>
<td>(2meptH3)4PbBr4</td>
<td>(0.24, 0.23)</td>
<td>91</td>
<td>-</td>
<td>2.23 (av)</td>
<td>3.37</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td>(EDBE)2PbBr4</td>
<td>(0.39, 0.42)</td>
<td>84</td>
<td>5509</td>
<td>14 (λem = 570 nm)</td>
<td>9.0</td>
<td>7</td>
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<tr>
<td>&lt;110&gt; 2D perovskite</td>
<td>(N-MEDA)[PbBr4]</td>
<td>(0.36, 0.41)</td>
<td>82</td>
<td>4669</td>
<td>1.2</td>
<td>1.5</td>
<td>55</td>
</tr>
<tr>
<td></td>
<td>(AP)2PbBr4</td>
<td>(0.24, 0.32)*</td>
<td>65*</td>
<td>11702*</td>
<td>-</td>
<td>-</td>
<td>21</td>
</tr>
<tr>
<td></td>
<td>(AETU)2PbBr4</td>
<td>(0.16, 0.18)*</td>
<td>-</td>
<td>&gt;100000*</td>
<td>-</td>
<td>-</td>
<td>54</td>
</tr>
<tr>
<td>3X3 &lt;110–2D perovskite</td>
<td>α-(DMEN)2PbBr4</td>
<td>(0.28, 0.36)</td>
<td>73</td>
<td>7843</td>
<td>1.39 (λem = 545 nm; av)</td>
<td>-</td>
<td>24</td>
</tr>
<tr>
<td>cationic 2D lead halide</td>
<td>[Pb2F5][O2C(CH2)3CO2]-</td>
<td>(0.33, 0.41)</td>
<td>78</td>
<td>5620</td>
<td>2.64 (λem = 541 nm; av)</td>
<td>1.8</td>
<td>49</td>
</tr>
<tr>
<td></td>
<td>[Pb2Cl5][O2C(CH2)3CO2]-</td>
<td>(0.35, 0.46)</td>
<td>68</td>
<td>5129</td>
<td>2.24 (λem = 536 nm; av)</td>
<td>11.8</td>
<td>49</td>
</tr>
<tr>
<td></td>
<td>[Pb2Br5][O2C(CH2)3CO2]-</td>
<td>(0.42, 0.48)</td>
<td>-</td>
<td>3770</td>
<td>2.25 (λem = 565 nm; av)</td>
<td>2.0</td>
<td>49</td>
</tr>
<tr>
<td>1D nanotube</td>
<td>(HMTA)4PtBr8</td>
<td>(0.42, 0.45)</td>
<td>78*</td>
<td>3572*</td>
<td>106 (λem = 580 nm; av)</td>
<td>7%</td>
<td>61</td>
</tr>
<tr>
<td>1D perovskite</td>
<td>(DMEDA)4PbBr8</td>
<td>(0.21, 0.28)</td>
<td>53</td>
<td>21242</td>
<td>26 – 37 (λem = 475 nm; av)</td>
<td>18-20</td>
<td>40</td>
</tr>
<tr>
<td>0D perovskite</td>
<td>(DMEDA)2SnBr6</td>
<td>(0.45, 0.51)*</td>
<td>54*</td>
<td>3506*</td>
<td>2200 (λem = 570 nm; av)</td>
<td>95</td>
<td>63</td>
</tr>
<tr>
<td></td>
<td>(DMEDA)2SnI6</td>
<td>(0.58, 0.41)*</td>
<td>71*</td>
<td>1607*</td>
<td>1100 (λem = 620 nm; av)</td>
<td>75</td>
<td>63</td>
</tr>
<tr>
<td></td>
<td>(BMPY)2SnBr6</td>
<td>(0.50, 0.47)*</td>
<td>61*</td>
<td>2588*</td>
<td>4200 (λem = 590 nm; av)</td>
<td>98</td>
<td>63</td>
</tr>
</tbody>
</table>

Figure 7. CIE chromaticity coordinates of broadband emitting metal halide frameworks.

4 Applications of broadband polaronic emitters

The structural properties of low-dimensional metal halide perovskites can be synthetically engineered to modulate the PL energy and bandwidth, making them highly attractive for solid-state lighting (SSL) and integration in light emitting diodes and transistors. Table 1 and Fig. 7 summarize the luminescence properties for different types of broadband emitting perovskites in terms of color rendering index (CRI), correlated colour temperature (CCT), Commission Internationale de l’Eclairage (CIE) chromaticity coordinates and PLQY. In particular, the achievement of high CRI values (>80) is required for human eye-friendly and color-critical high-level applications and is crucial to make SSL competitive against conventional lighting and display systems. Nowadays, white-light emission in solid state devices is typically achieved by mixing different phosphors, leading to issues regarding the poor color rendition, self-absorption and different degradation rates. Broadband emitting perovskites are ideal candidates for application in single emitter white-light LEDs, helping to avoid the...
aforementioned issues. Although excellent CRI values were achieved with 3D haloplumbate frameworks and 2D perovskites, their PLQY is still low (<9%) for practical applications. Among layered perovskites, the <100> type with strong octahedral distortion provides the highest CRI values, although the PLQY are generally low. This suggests that the strong structural distortion necessary to favor exciton self-trapping in 2D perovskites also open additional non-radiative relaxation pathways partially quenching the luminescence. Increased quantum confinement effects in lower dimensional structures ease the formation of self-trapping phenomena and substantially increases the emission efficiency and PL lifetime, but tends to red-shift the luminescence (Table 1 and Fig. 7), worsening the quality of the white-light emission. To date, (EDBE)PbBr₄ microcrystals, (DMEDA)SnBr₆, (PEA)₂PbBrClₓ and Cs₂Ag₀.6Na₀.4InCl₆ have been employed for fabrication of white-light emitting LEDs using a down-conversion approach, where a UV commercial LED was used to excite the perovskite.⁶³,⁶⁴,¹¹⁴ As alternative approach for the emission tunability, the ultrabroad PL spectrum of these materials could be modified by engineering their photonic environment so that the PL is suppressed or enhanced at specific spectral regions. In this regard, we have recently shown the directional spectral redistribution of (EDBE)PbCl₄ luminescence by incorporation in a polymer microcavity (Fig. 8a).¹¹⁵ Similarly, perovskite films could be patterned on the sub-wavelength scale (for example through large area techniques like nanoimprint lithography) to enhance the density of photon states leading to Purcell enhancement of luminescence and structural coloring.¹¹⁶ In future, these strategies could be applied to allow color tunability in low-cost, flexible LEDs based on an unique broadband emitter without relying on compositional engineering.¹¹⁵

Figure 8. a) Distributed Bragg reflector (DBR) coupled to the perovskite [EDBE]PbCl₄, PS = polystyrene and CA = cellulose acetate; Reprinted with permission from ref. 115 Copyright 2018 American Chemical Society. b) X-ray excited luminescence of (EDBE)PbCl₄ reprinted with permission from ref. 10 Copyright 2016 Springer Nature.

Thanks to the solution processability and the presence of heavy atoms which enhance their absorption cross section, hybrid perovskites are also promising for the fabrication of cost-effective, large area X-ray detectors,¹¹⁷-¹¹⁹ which find application in scientific research as well as medicine and inspection systems for safety controls. Due to the high exciton binding energy and large Stokes-shift, which reduce the negative effect of thermal quenching and the probability of radiation self-absorption, we have shown that X-ray scintillators based on the white-light emitting (EDBE)PbCl₄ outperform those based on MAPbI₃ and MAPbBr₃. In all these perovskites, following X-ray excitation, an afterglow effect extending to thousands of seconds was observed at low temperature (10 K), while showing a negligible contribution at room temperature. The fraction of the total excitation energy accumulated into long-living traps was determined by thermoluminescence measurements, which allows to compare the luminescence intensity from the end of X-ray excitation until the end of the entire run (Iₑ), against the luminescence intensity from the beginning of the X-ray excitation to the end of the run (Iₑ + Iₑₑ). The ratio Iₑ/Iₑₑ was found to be ~0.002 for 3D perovskite and ~0.058 for (EDBE)PbCl₄, which are both lower compared to other oxide-based scintillator materials.¹⁰ In particular, higher luminescence yields were achieved in the range 50-298 K (Fig. 8b) with the white emitter (EDBE)PbCl₄ as compared to the standard 3D perovskites, reaching 9,000 photons/MeV at room temperature.¹⁰ Such encouraging performance might be further improved by increasing the luminescence quantum yield of the 2D perovskite through structural design, by doping with lanthanide ions such as Ce³⁺ or by enhancing the X-ray luminescence yield using nanostructured perovskite films (for example via Purcell enhancement).

Beyond the efficient light emitting properties, metal halide perovskites are also emerging for their attractive gain characteristics,¹²⁰-¹²³ which recently allowed to achieve continuous-wave (c. w.) optically pumped lasing (at T < 160 K) in distributed feedback lasers based on the 3D perovskite MAPbI₃.¹²⁴ Further improvements are now being achieved with layered perovskites, in particular multidimensional perovskites of the Ruddlesden-Popper series, where energy funneling effects in heterogeneous systems allowed to obtain low-threshold lasing (13.6 µJ/cm²) and improved gain coefficients (> 300 cm⁻¹).¹²⁵-¹²⁷ Such 2D/3D systems can potentially be engineered to obtain an optimal balance between efficient radiative recombination, long radiative lifetime and high carriers mobility,¹²⁸ and might hold the key for the realization of the long sought-after solution processed laser diodes.¹²⁷ In this context, white light emitting perovskites might offer the attractive possibility to realize tunable lasers across the visible spectral region with a single active material. Nevertheless, the capability to sustain amplified spontaneous emission and lasing in such broadband
emitters is yet to be ascertained, urging the need for a deeper understanding of the photophysical processes underlying the broadband luminescence of metal halide lattices.

To further widen the application prospective of broadband emitting perovskites (including, for example, light-emitting field effect transistors and electrically injected lasers), future works will also need to address charge transport and injection issues in these wide band-gap semiconductors as well as elucidate the non-trivial role of small-polaron transport in metal halide frameworks. Indeed, while electroluminescence using narrowband-emitting 2D, multidimensional and 3D perovskites as active materials is now established, the direct electrical to optical power conversion in white-light perovskites still remains a challenge.

5 Conclusions

Recent progress in synthetic design suggests that ultrabroadband and largely Stokes shifted luminescence is an intrinsic characteristic of a wide range of metal halide frameworks with low dimensionality. Up to now, the most accredited model to explain this behavior points towards the formation of several self-trapped excitonic species, which can be interpreted as small polarons localized at specific sites of the inorganic lattice. The balance between free and self-trapped excitons is strongly affected by temperature and by the local structural properties of the materials, where stronger deviations from the ideal geometry seem to ease the self-trapping process. While this has been more extensively investigated in 2D perovskites, the analysis of a broader range of low-dimensional structures will be necessary to univocally define the common structural properties which stabilize the polaron formation. Several studies are still needed to provide deeper insights into the trapping mechanism, such as electron paramagnetic resonance, neutron diffraction, infrared photoinduced absorption and time resolved photocurrent. These will provide additional information to correlate structure and optical properties to aid the design of highly efficient white-light emitters for optoelectronic applications, such as light emitting devices and X-ray scintillators.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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Notes and references


32. I. Neogi, A. Bruno, D. Balahayian, T. W. Goh, B. Ghosh, R.

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Structural properties closely affect the photophysical processes underlying photoluminescence broadening and white-light emission in low-dimensional perovskites.