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3, 7061Emission colours of bistable photochromic
compounds: Ce-doped alkaline (Rb, K, Na)–
indium fluorides†Manuel Gaudon,^{*a} Ines Andron,^a Alain Demourgues,^a Vincent Rodriguez,^{id b}
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The structural and spectroscopic properties of three Ce³⁺-doped A₃InF₆ compounds (with A⁺ = Rb⁺, K⁺, Na⁺) are investigated. Containing indium, they all exhibit 2Ce³⁺ + In³⁺ ⇌ In⁺ + 2Ce⁴⁺ redox processes induced by UV irradiation between Ce³⁺ and In³⁺ cations. The photochromo-luminescent behaviour and associated kinetics, with the Ce³⁺ and In⁺ ion emission quenching through the redox equation, respectively, are investigated. Our studies highlight that different mechanisms (surface vs. bulk) need be considered to explain the decrease in the blue (Ce³⁺) and orange (In⁺) emission curves. Moreover, the comparison of Ce³⁺-doped Rb₂KInF₆, K₂NalInF₆ elpasolites and K₃InF₆ cryolite spectral distributions shows that changing the nature of the alkali cation makes colour tuning possible. The shift of the Ce³⁺ and In⁺ colour emissions between the three as-prepared compounds is mainly due to the variation in the ionic-covalent nature of the chemical bonds around the active cation sites. The final calculated optical difference between the two bistable states varies from 0.31 to 0.48 depending on the alkali subnetwork.

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1. Introduction

As photochromism can be defined as the ability of a compound to change its colouration under UV or visible irradiation [...], the photochromoluminescence property is herein defined as the ability of the studied inorganic fluorides to reversibly change their emissive colouration resulting from UV irradiation. Hence, in photochromoluminescent materials, the A colour emission state is shifted to the B colour emissive state from irradiation at a λ_A wavelength (forth-irradiation), whereas reversibly, the B state can return to the A state from irradiation at a λ_B wavelength (back-irradiation). Such compounds can be developed in view of various applications, such as information storage,^{1–3} UV sensors,⁴ and security inks.⁵

The studied compounds are inorganic indium-based fluorides, with a double perovskite-type structure, constituted mainly on alkali–fluorine ionic bonds with the general formula A₂BKInF₆ (A and B being alkaline elements).⁶ Compounds were doped with trivalent cerium in substitution of trivalent indium, with a substitution concentration of 2 mol%. Both cerium and indium ions are present at the source of the potential modulation in the emission of the Ce-doped fluorides. Indeed, in such inorganic fluorides: RE-doped A₂BKInF₆ series (RE: rare earth element), a redox process can be produced between cerium and indium cations, as demonstrated by Chaminade *et al.*,⁷ or more recently developed by Cornu *et al.*^{8,9} It was shown in the last former article that the UV irradiation of Ce³⁺-doped Rb₂KInF₆ elpasolite-type compounds induces a reversible redox process.

Under UV irradiation at 315 nm, the progressive disappearance of two Ce³⁺ ions in favour of the appearance of one In⁺ cation is associated with the Ce³⁺ blue emission quenching and the increase in the orange In⁺ emission (forth-irradiation). A reversible phenomenon is observed under material irradiation at 255 nm (back-irradiation). A nice on–off effect of the Ce³⁺/In⁺ cation emission is obtained; the cycling of this redox process has been checked over 5 cycles under UV irradiation; and the stability of both Ce³⁺–In³⁺ (A state) and Ce⁴⁺–In⁺ (B state) states was confirmed over 10 years.⁹

In this paper, attempts to observe a redox phenomenon similar to that in Ce-doped Rb₂KInF₆ compounds have been

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performed on the structurally close compounds: Ce-doped K_2NaInF_6 elpasolite and Ce-doped K_3InF_6 cryolite. One target is to tune the colour contrast between the two states (forth- and back-irradiation) in this series. The studied halide compounds are all related to the perovskite structure. The latter are generally studied for their wide range of physical properties, such as ferroelectricity, dielectricity, pyroelectricity and piezoelectricity.¹⁰ The crystallographic structure of the three as-prepared compounds (Ce-doped Rb_2KInF_6 was again prepared as a reference) was studied by X-ray diffraction and Raman spectroscopy in the first part. In the second part, the luminescence properties of the cerium or indium luminescent centres in these three compounds are compared and interpreted based mainly on the Stokes shift consideration.¹¹ The kinetic mechanism of Ce^{3+} luminescence quenching (forth redox) and In^+ luminescence (back redox) quenching is deeply investigated.

2. Experimental

The elpasolite/cryolite compounds were prepared from the solid-state route in sealed tubes.

$\alpha\text{-InF}_3$ was issued from direct fluorination using diluted F_2 (10%, Ar filled) gas treatment at $T = 450^\circ\text{C}$ during 6 hours of $\text{InF}_3 \cdot x\text{H}_2\text{O}$, hydrated fluorides, which were prepared from a coprecipitation process.

The reagents RbF , KF , NaF , CeF_3 fluorides (commercial compounds, Aldrich) and InF_3 used for the synthesis were hygroscopic, so they were previously treated at 110°C for 3 hours under vacuum to remove all traces of water and then placed in a glove box (BAG). The reagents were then weighed in stoichiometric proportions and ground in an agate mortar. They were introduced into a clamped argon atmosphere glove box and sealed platinum tube to prevent any air entry. The tube was placed in a muffle furnace and heat-treated at 700°C for 12 hours.

Powder X-ray diffraction patterns were collected on a Philips X'Pert MPD X-ray diffractometer with a Bragg–Brentano geometry using $\text{Cu K}\alpha_{1,2}$ radiation ($10^\circ < 2\theta < 130^\circ$, step 0.02° and counting time of 30 seconds). The diffractograms were refined using the Rietveld refinement method with the conventional reliability factors for refinement quality assessment. The Fullprof program package was used. Unit-cell parameters, zero-shift, peak profile parameters, atomic positions and isotropic displacement factors were refined; however, occupancies were fixed equal to the target composition.

Raman spectra were recorded at room temperature on a Xplora confocal Raman microscope from Horiba Scientific (objective $50\times$, $\text{NA} = 0.50$) using a CW 532 nm laser excitation. The laser power was less than 15 mW for all samples. The spectrometer includes a grating with 2400 grooves per mm and an air-cooled CCD camera for detection. Unpolarized Raman spectra were measured over the frequency range $150\text{--}750\text{ cm}^{-1}$ with a spectral resolution better than 3 cm^{-1} . The spectral range below 150 cm^{-1} has been deliberately rejected because the longpass edge filter for the collection of the Raman Stokes

signal strongly disturbs the profile and the intensities of the peaks possibly present in this range.

The photoluminescent properties were analysed using a Fluorolog 3 Horiba spectrofluorimeter connected by a 450 W continuous Xenon source. Raw data were corrected for the experimental parameters: variation of the incident flux, monochromator transmission and photomultiplier sensitivity.

3. Results and discussion

3.1 Structural investigation

Perovskite designates any ABX_3 structural material, formed on the basis of a three-dimensional network of corner-sharing BX_6 octahedra in all three spatial directions and with large A cations occupying all cuboctahedral cavities, delimited by the eight BX_6 octahedra (Fig. 1).¹² The studied halogenated perovskites have the generic formula $\text{A}_2\text{BB}'\text{F}_6$ with a strict alternation of the B (Na^+ or K^+) and B' (In^{3+}) cations that causes the doubling of the lattice parameter (ordered double perovskites). The Rb_2KInF_6 (labelled Rb_2K) and K_2NaInF_6 (labelled K_2Na) compounds both crystallize in this so-called elpasolite structure. Finally, a special case of elpasolite is the A_3BX_6 cryolite (named after the Na_3AlF_6 archetypal mineral), in which the A atoms occupy half of the octahedra and all cuboctahedron sites, as for the K_3InF_6 (labelled K_3) compound.

The X-ray diffraction patterns of the three studied compounds are depicted in Fig. 2. The as-quoted Rb_2K and K_2Na compounds both crystallize in the $Fm\bar{3}m$ space group, as already attested by the literature (ICSD files 00-048-1754 and 00-023-1361,

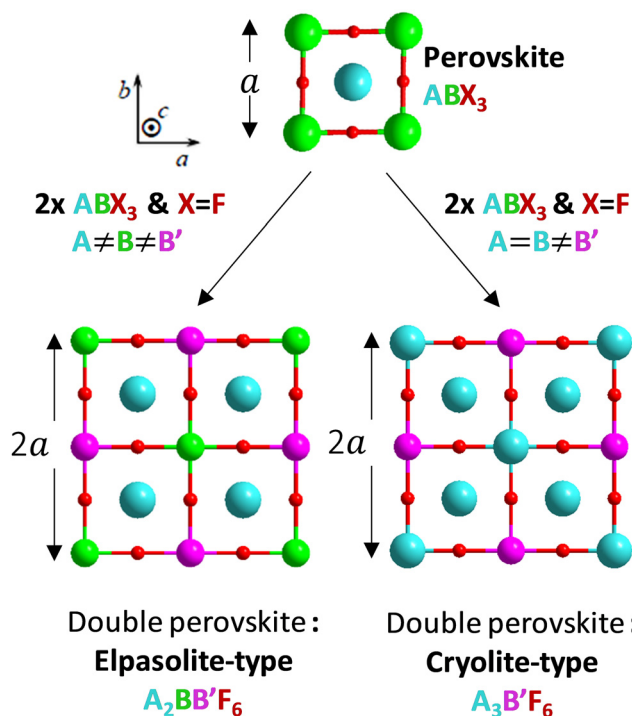


Fig. 1 Structural filiations linking the perovskite framework and the two studied double perovskite types: elpasolite-type and cryolite-type.



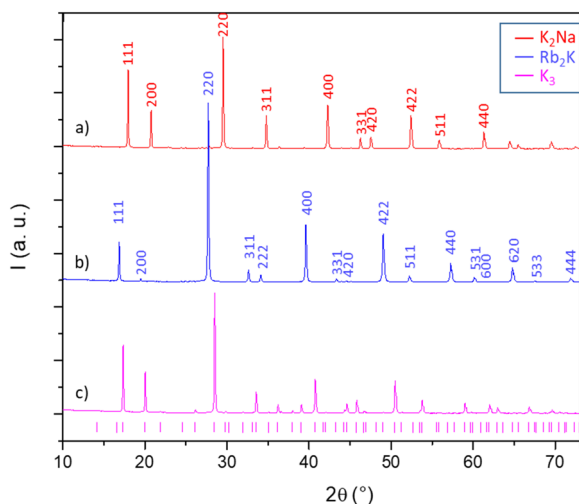


Fig. 2 X-Ray diffraction patterns of the three studied compounds: (a) K_2Na , (b) Rb_2K (both with $Fm\bar{3}m$ space group) and (c) K_3 (with intermediate peak indexation corresponding to $Fd\bar{3}$ space-group for the latter).

respectively, for Rb_2K and K_2Na). Moreover, the clear right-shift of the peak positions of the K_2Na phase with respect to the Rb_2K phase already shows that the unit-cell parameter of the first is the smallest. The K_3 compound has a peak series that seems to coincide with the 01-072-1763 file ($Fd\bar{3}$ space group), but the peak assignment is not fully satisfying and deserves deeper investigation.

In the $A_2BB'F_6$ investigated compositions, the ionic radius into octahedral sites, either the B ions (1.38 Å, 1.02 Å, for K^+ and Na^+ , respectively) or the B' ions (0.8 Å for In^{3+}), are too high compared to the radii of A cations, which are located in cuboctahedral sites (1.72 Å and 1.64 Å for Rb^+ and K^+ , respectively). Both the compound frameworks are subjected to internal stresses associated with a Goldschmidt tolerance factor, less than 1, causing the tilting of the octahedra chains to satisfy the valence of the whole cation set.¹³ As previously shown in our former study devoted to Rb_2K compound,^{8,9} considering a $Fm\bar{3}m$ cubic system for the first two compounds (K_2Na , Rb_2K), the octahedra rotation results in a positioning of the fluorine ions in the 96j Wyckoff

position, with an equiprobable random occupancy of a quarter compared to the 24e position located on the In–B–In axes. The 96j fluorine position is distinguished from the 24e position by off-centering the In–B–In axes by a distance that is characterized by the fluorine z coordinate.

The K_3 compound has a Goldschmidt factor significantly lower than 1 ($t = 0.868$), and therefore, a lower symmetry than the $Fm\bar{3}m$ cubic cell is expected.¹⁴ From Glazer theory and the fixed symmetry relations between space groups, we assumed that the K_3 phase could crystallize in a cell with an $I4/m$ space group. Furthermore, this space group corresponds to the low-temperature polymorph of Rb_2K .¹⁵ Refinements were performed maintaining a “doubled” c parameter with respect to the $Fm\bar{3}m$ elpasolite unit-cell. However, the refinement leads to low reliability factors. Thus, a full resolution of the K_3 crystalline framework is proposed with the $I4/m$ space group. There are four positions for indium atoms and three positions for potassium atoms. The twelve-coordinated potassium atoms are located in two different 16i positions. Finally, there are ten positions for fluorine atoms. The indium and potassium octahedra constituting the $I4/m$ double-cell of the K_3 compound are depicted in Fig. 3, and each distinct octahedron (peculiar Wyckoff position) is represented by a different edge colour.

Table 1 summarizes the refined parameters and reliability factors of the structural hypotheses carried out for fluorine ions at position 24e or 96j on Rb_2K and K_2Na , considering the just previously defined $I4/m$ double-cell for K_3 (detailed atomic positions for this latter compound are reported in Table S1a, ESI†).

Obviously, the unit-cell volume per unit formula (Z) is directly related to the alkaline ionic size of the compounds; thus, the equivalent unit-cell volume ($Z = 4$) decreases from Rb_2K ($V = 751.6 \text{ Å}^3$, $Z = 4$), then K_3 ($V/4 = 692.47 \text{ Å}^3$, $Z = 16$), and finally to K_2Na ($V = 623.95 \text{ Å}^3$, $Z = 4$).

For the first two compounds (Rb_2K and K_2Na), the positioning of fluorine in the 96j position instead of the 24e position makes it possible to significantly reduce the reliability factor values for the two systems. We also observe a decrease in the B_{iso} isotropic displacement factors associated with fluorine ions. The fluorine

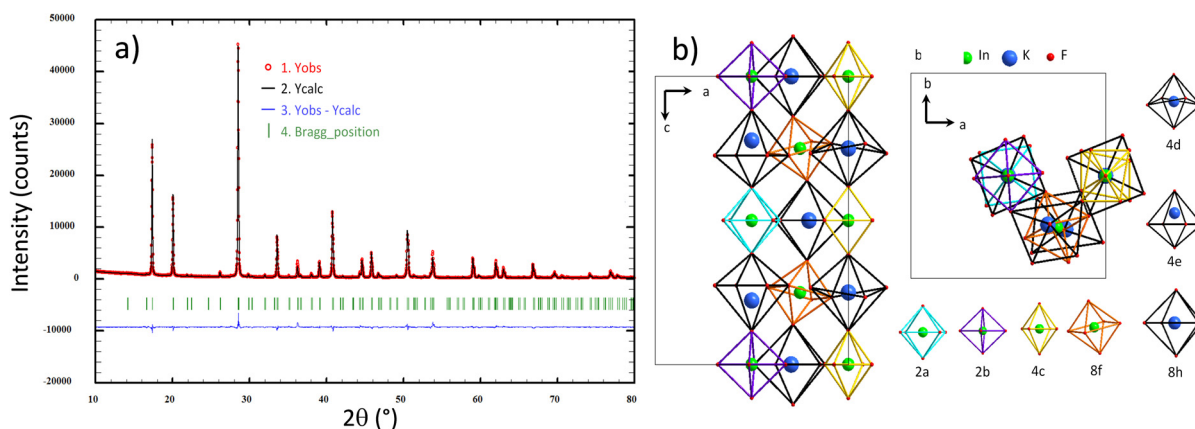


Fig. 3 Rietveld refinement considering the $I4/m$ space group for the K_3 compound (a) and representation of all B and B' cationic sites and their crystallographic subframework for the K_3 compound (b).



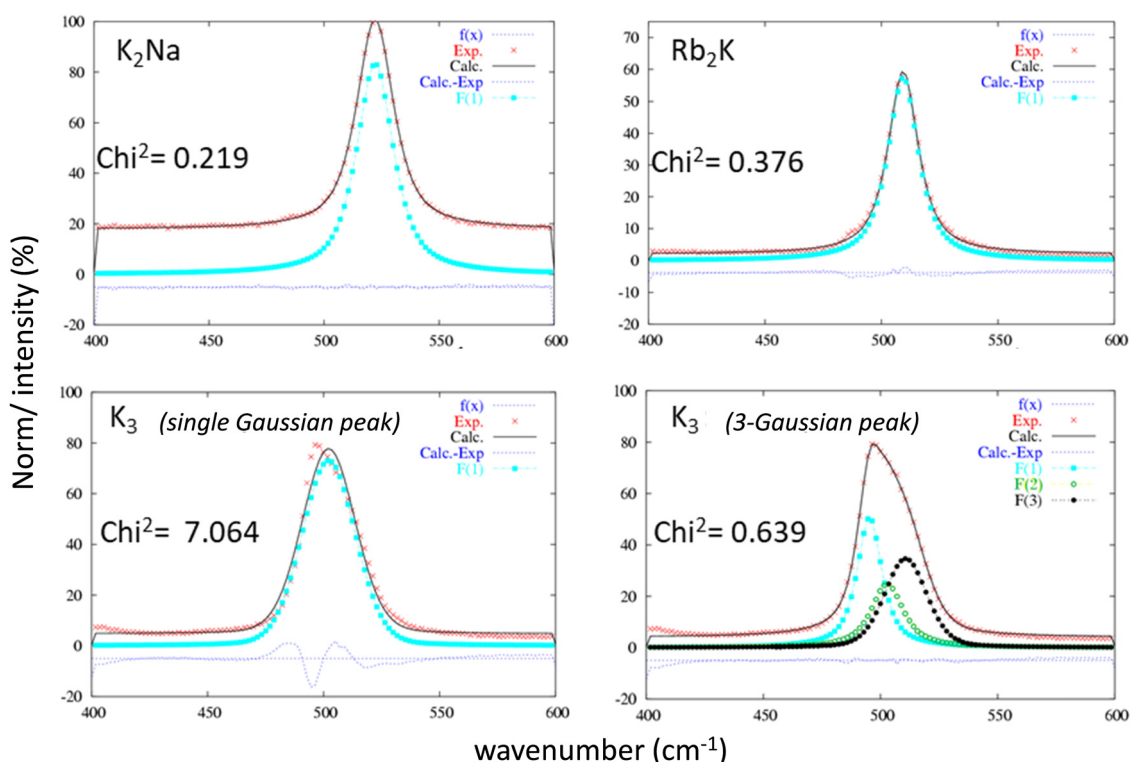
Table 1 Unit cell parameters, fluorine coordinates and reliability factors extracted from the X-ray powder pattern refinement of the three studied compounds

F atom position	Rb ₂ K – <i>Fm</i> $\bar{3}$ <i>m</i>		K ₂ Na – <i>Fm</i> $\bar{3}$ <i>m</i>		K ₃ – <i>I</i> 4/ <i>m</i>
	24c	96j	24c	96j	
<i>a</i> (Å)	9.0922(3)	9.0921(2)	8.5468(2)	8.5450(3)	(<i>a</i> $\sqrt{2}$) 12.5095(3)
<i>c</i> (Å)	—	—	—	—	(2 <i>a</i>) 17.6890(7)
<i>x</i> (F)	0.2317(4)	0.2317(3)	0.2438(4)	0.245(4)	Detailed atomic positions – see Table S1 (ESI)
<i>z</i> (F)	0	0.041(1)	0	0.045(1)	
<i>V</i> (Å ³)	751.64(3)	751.60(4)	624.33(3)	623.95(3)	2769.9(3) (<i>Z</i> = 16)
<i>B</i> _{iso} (A) (Å ²)	2.70(5)	2.84(5)	1.73(6)	1.75(6)	0.83(4) all cations together
<i>B</i> _{iso} (B) (Å ²)	1.3(10)	1.27(9)	1.3(1)	0.86(2)	
<i>B</i> _{iso} (B') (Å ²)	0.89(4)	0.97(4)	0.39(4)	0.54(3)	
<i>B</i> _{iso} (F) (Å ²)	7.2(1)	3.3(2)	4.8(1)	1.3(2)	1.0(2)
<i>R</i> _p	11	10.8	14.1	13.4	10.1
<i>R</i> _{wp}	11.3	10.8	15.2	14.4	11.2
Bragg <i>R</i> -factor	5.35	3.43	4.12	2.98	4.27
χ^2	7.05	4.91	8.74	7.83	4.62
Scorr	3.5	3.4	2.9	2.9	3.2
Goldschmidt (<i>t</i>)	0.891		0.938		0.868

atom is closer to the position $x = 0.25$ (midpoint between the B and the B' cation) for the K₂Na phase than for Rb₂K.

For the K₃ sample, a very complex framework based on multiple octahedral sites was evidenced. There are indeed 3 kinds of octahedron sequences along the *c* axis and 3 successive layers of atoms in the (*ab*) plane (Fig. 3b). Along the *c* axis, the potassium and indium octahedra generally tilt in the opposite direction from each other, as also observed for the neighbouring octahedra within each layer.

Raman spectroscopy studies of the InF₆ entities were undertaken to complete the X-ray diffraction studies. This optical spectroscopy is particularly sensitive to bond vibrations and allows the instantaneous environment of each ion to be probed, including dynamic distortions at short time scales down to a few tens of femtoseconds (fs). As a reminder, both the Rb₂K and K₂Na phases exhibit a single Wyckoff position for indium and potassium ions. The InF₆ octahedral sites are perfectly regular.¹⁶ In contrast, according to previous X-ray diffraction

**Fig. 4** Refinement of the ν_5 (F_{2g}) vibration band attributed to each of the three studied compounds considering a single Gaussian peak or a 3-Gaussian peaks for the K₃ compound.

studies, for the K_3 compound, the different octahedral sites show strong disparities in their geometry. In the three samples, the InF_6 octahedral sites show 3 active Raman modes: ν_1 (A_{1g}), ν_2 (E_g) and ν_5 (F_{2g}). The doubly degenerated antisymmetric valence vibration ν_2 (E_g) has a very low intensity at approximately 380 cm^{-1} . Therefore, we have concentrated our analysis on the two other vibration modes: they correspond to the three degenerate ν_5 (F_{2g}) shear modes and the nondegenerate ν_1 (A_{1g}) breathing mode. Whether for the ν_5 (F_{2g}) (Fig. 4) or ν_1 (A_{1g}) (Fig. 5) vibration mode, the same observations leading to the same conclusions can be formulated. For these two vibration modes, the Raman intensity profiles of the compounds $K_2\text{Na}$ and Rb_2K are perfectly refined by considering that the vibration band comes from a single Gaussian contribution. However, the two vibration signals of the K_3 compound fit better by considering the band resulting from the convolution of at least three Gaussian contributions (the χ^2 value is drastically decreased from 7.064 to 0.639, while 3 Gaussian contributions are taken into account for the fitting). This Raman analysis therefore clearly reflects the great disparity of In sites and In-F distances in cryolite-type structures.

The parameters extracted from the refinement of the ν_5 (F_{2g}) and ν_1 (A_{1g}) bands for the three studied compounds are listed in Table 2. Two additional observations can be made, considering that the frequency of vibration mode is higher than this one: (i) the respiration mode is strongly constrained by the presence of sodium Na^+ instead of K^+ on site B (position at 522 cm^{-1} for $K_2\text{Na}$, approximately $502\text{--}509\text{ cm}^{-1}$ for Rb_2K and K_3); (ii) the shear mode is strongly relaxed by the presence of Rb^+

Table 2 Parameters extracted from the refinement of the ν_5 (F_{2g}) and ν_1 (A_{1g}) bands

		$K_2\text{Na}$	Rb_2K	K_3	K_3	
	Refined parameters	1-Gaussian envelope			3-Gaussian envelope	
ν_5 (F_{2g})	$d(\text{In-F})_{\text{moy}}$ (\AA)	2.12(1)	2.14(1)	2.13(1)	2.13(1)	
	Position (cm^{-1})	233	218	230	218	230
	Width (cm^{-1})	14	16	27	26	15
ν_1 (A_{1g})	Intensity (a.u.)	1.2	0.9	1.1	0.7	1.1
	Position (cm^{-1})	522	509	502	495	502
	Width (cm^{-1})	19	16	27	13	16
	Intensity (a.u.)	1.2	0.9	1.1	1.3	0.7

ions instead of K^+ on site A (218 cm^{-1} for Rb_2K instead of $230\text{--}233\text{ cm}^{-1}$ for K_3 and $K_2\text{Na}$).

3.2 Luminescence properties: photochromoluminescence.

Doped with 2% (molar substitution at the indium site) cerium ions, Rb_2K , $K_2\text{Na}$ elpasolites and K_3 cryolite present $4f \rightarrow 5d$ electronic transitions under UV irradiation. Fig. 6 shows the corresponding excitation and emission graphs normalized at the maximum. Table 3 lists the Stokes shift values calculated as the wavenumber difference (cm^{-1}) between the maximal intensity of the lowest $5d(T_{2g})$ excitation level and the corresponding $4f(^2F_{5/2})$ arrival level. The emission band is clearly constituted from the overlapping of two Gaussian contributions, indexed to the $5d(T_{2g}) \rightarrow 4f(^2F_{5/2})$ and $5d(T_{2g}) \rightarrow 4f(^2F_{7/2})$ transitions. The deconvolution of the K_3 emission is reported as an illustration (Fig. 6c). Energy differences of 2897 cm^{-1} , 2347 cm^{-1} and

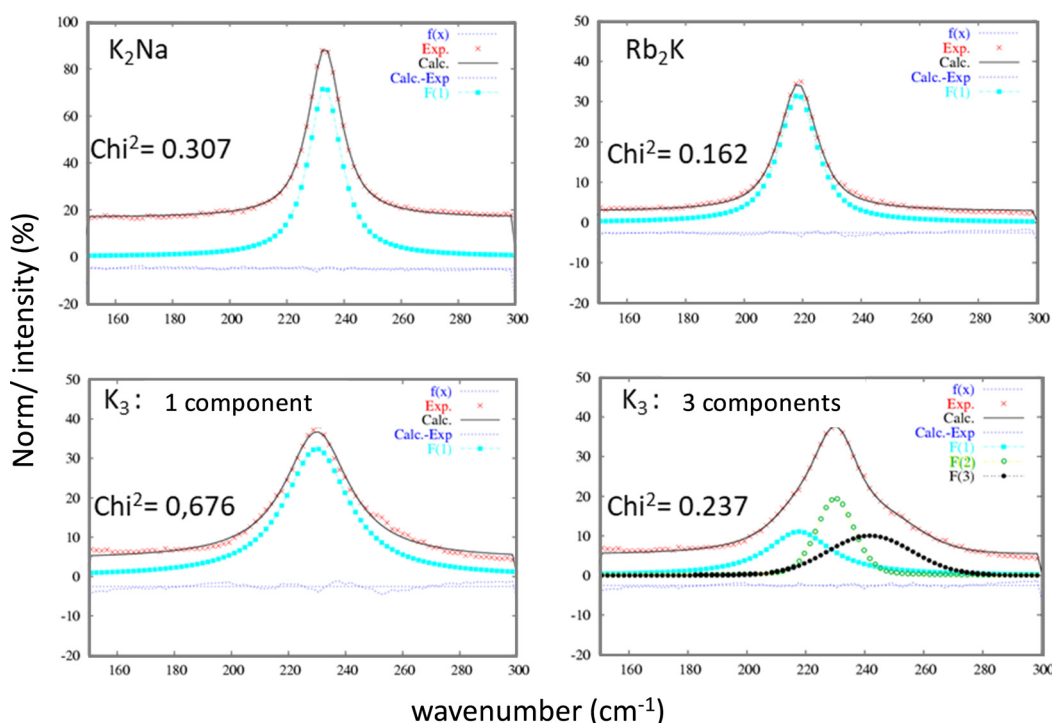


Fig. 5 Refinement of the ν_1 (A_{1g}) vibration band attributed to each of the three studied compounds considering a single Gaussian peak or a 3-Gaussian peak for the K_3 compound.



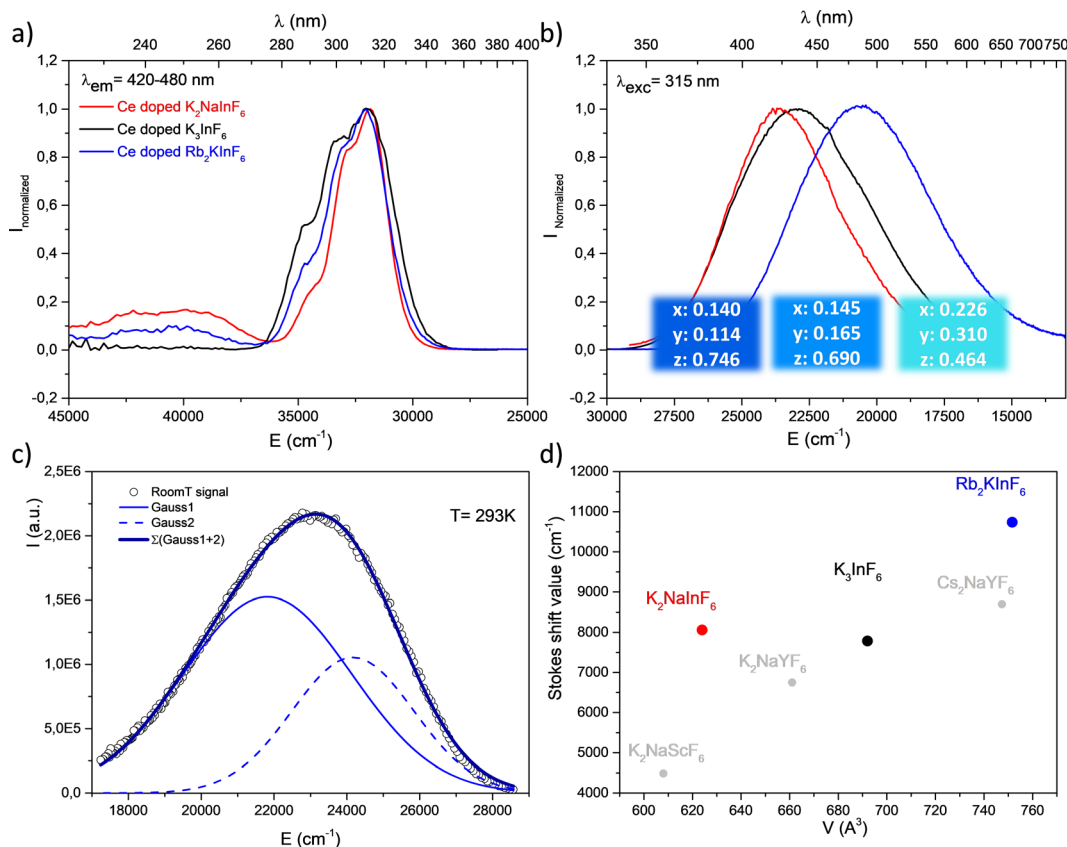


Fig. 6 Excitation spectra (a), emission spectra and corresponding colour coordinates (b), deconvolution of the emission spectra into two $5d(T_{2g}) \rightarrow {}^2F_{7/2}$ and $5d(T_{2g}) \rightarrow {}^2F_{5/2}$ bands at room temperature of K_3 (c) and Stokes shift of Ce^{3+} doped-fluoride compounds (d).

2424 cm^{-1} for Rb_2K , K_2Na and K_3 , respectively, are observed, in agreement with the expected value resulting from the splitting of the two Ce^{3+} 4f levels by spin orbit coupling.¹⁷ Regardless of the nature of the host lattice, the maximum of the excitation band is located at approximately 313 nm, which indicates a very weak difference in the crystal field effect on the 5d orbital splitting. However, the global emission is significantly shifted to the highest energy *versus* the decrease in the alkali size (as a reminder, $r_{Rb}[6/12] = 1.52/1.72$ Å; $r_K[6/12] = 1.38/1.64$ Å; $r_{Na}[6/12] = 1.02/1.39$ Å). Thus, the Rb_2K and K_2Na emission bands peak at 485 nm and 423 nm, respectively. The K_3 emission is maximum at 435 nm. The corresponding trichromatic x , y , z coordinates associated with luminescence colouration are indicated in Fig. 6b (colour panels are also proposed due to the conversion of xyz parameters into the RGB system).

Stokes shift values (Δ_{ss}) follow the order $\Delta_{ss} Rb_2K > \Delta_{ss} K_2Na > \Delta_{ss} K_3$. The standard deviation of the In–F distances at the Ce^{3+} site is equal to 0.01 Å (Table 2); this parameter cannot

explain such an evolution. Δ_{ss} reflects both the variation of the equivalent unit cell volumes ($Rb_2K > K_3 > K_2Na$) and the breathing of the InF_6 polyhedra. As discussed previously, a larger volume results in an increased disorder of the fluorine anions (see Table 1). In particular, considering that the softness of the structure increases with alkali size, we suggest that the capability of a larger Ce^{3+} coordination polyhedron to be distorted, leading to its energetic and spatial reorganization. Thus, the cryolite structure appears as a more rigid host lattice. Such a tendency seems to be confirmed by experimental data reported in the literature for similar structural matrices (Fig. 6d). Low-temperature measurements were performed on the K_3 compound (an equivalent study has been reported for the Rb_2K elpasolite.⁸ A weak thermal quenching of 12% (integrated area), a decrease in the full width at half maximum, and a UV shift are observed from a drop in the temperature down to 8 K. The results are illustrated in Fig. S1 (ESI†).

Table 3 Energy of the Ce^{3+} ion transitions in elpasolite (*cryolite) compounds

Compound	$V(Z=4)$ (Å ³)	λ_{ex} max T_{2g} (cm ⁻¹)	λ_{em} max (cm ⁻¹)	$T_{2g} \rightarrow {}^2F_{5/2}$ (cm ⁻¹)	$T_{2g} \rightarrow {}^2F_{7/2}$ (cm ⁻¹)	Δ_{ss} (cm ⁻¹)	ΔE (cm ⁻¹) ${}^2F_{5/2} \leftrightarrow {}^2F_{7/2}$
K_2Na	623.95(3)	31 949 (313 nm)	23 641 (423 nm)	23 891 (419 nm)	21 467 (466 nm)	8058	2424
* K_3	692.01(8)	31 949 cm ⁻¹ (313 nm)	22 989 (435 nm)	24 165 (414 nm)	21 818 (458 nm)	7784	2347
Rb_2K	751.60(4)	31 949 cm ⁻¹ (313 nm)	20 619 (485 nm)	21 212 (471 nm)	18 315 (546 nm)	10 737	2897

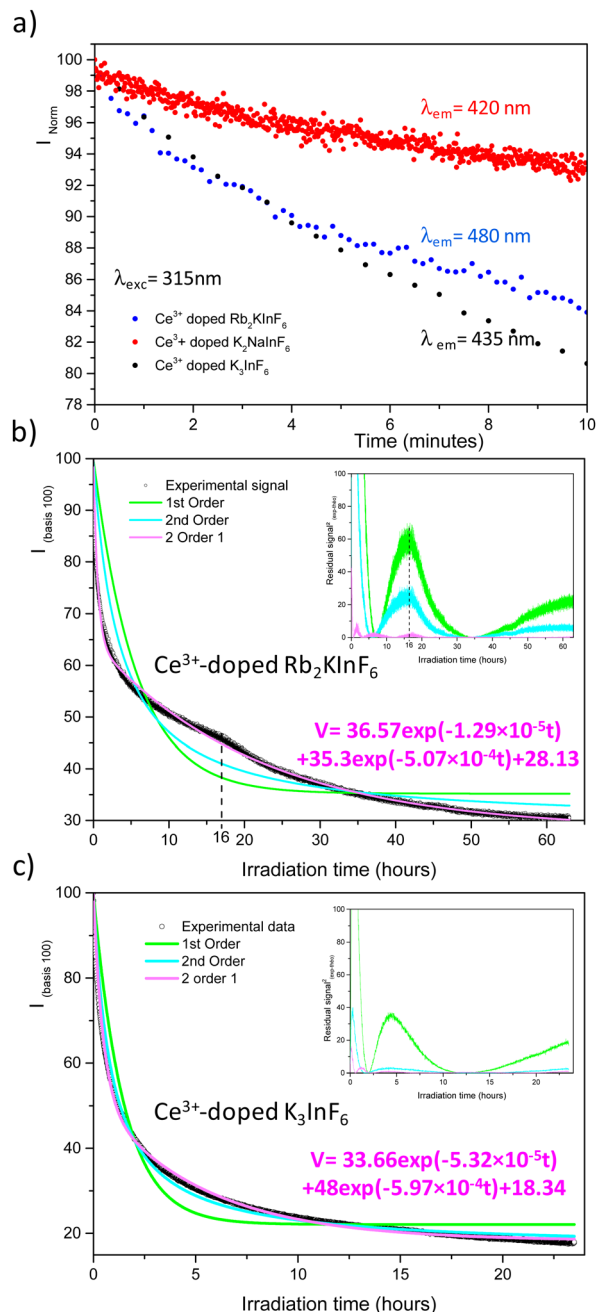


Fig. 7 Kinetic curves of Ce-doped Rb_2K and K_2Na elpasolites K_3 cryolite (a); fit of a long-time acquisition of the kinetic curves of Rb_2K and K_3 compounds. The square value of the residual difference between the calculated and experimental values is reported in inset (b and c). The fitting equations based on the iii model are indicated in pink.

As published previously, the emission of Ce^{3+} -doped Rb_2KInF_6 elpasolite is not stable under UV excitation. An electron transfer occurs between the rare earth and indium cations, leading to a redox process (fourth reaction) between the two elements. It is thus possible to record the kinetics of Ce^{3+} ion oxidation by following the decrease in the corresponding emission curve. As illustrated below (Fig. 7a), the redox process has also been detected on the K_2Na and K_3

compounds. After 10 minutes, losses in intensity of approximately 15%, 6% and 20% are observed for the Rb_2K , K_2Na and K_3 compounds, respectively. A deeper characterization is proposed for both the Rb_2K elpasolite and K_3 cryolite. The kinetic curves of the Ce^{3+} ions were recorded under excitation at 315 nm and a maximum emission fixed at 485 nm and 435 nm for Rb_2K and K_3 , respectively. On one hand, the oxidation velocity of Ce^{3+} cations and their disappearance have been considered equal to the global emission. Considering this hypothesis for the low doping rate, the emission intensity is proportional to the concentration of luminescent cations and can be expressed as $V = -d[\text{Ce}^{3+}]/dt$ avec $[\text{Ce}^{3+}]$ with $[\text{Ce}^{3+}]$, the concentration and t , the time. On the other hand, several kinetic models following the general rule $V = k[\text{Ce}^{3+}]^\alpha$ with α , the order of the reaction and k , the velocity constant, have been run by changing the α value.

The results of the fits are illustrated in Fig. 7b and c, and the corresponding parameters are listed in Table S2 (ESI[†]). The R^2 reliability factor was calculated as follows.^{18,19}

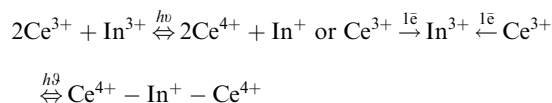
$$R^2 = 1 - \sqrt{\sum (x_{\text{signal}} - x_{\text{theo}})^2 / x_{\text{theo}}^2} \times 100,$$

where x_{signal} is the experimental value and x_{theo} is the simulated value.

Different hypotheses were considered. (i model) is a single first-order kinetic model corresponding to the disappearance of a single cerium luminescent centre such as $-d[\text{Ce}^{3+}]/dt = k_1[\text{Ce}^{3+}]^1$, leading to $[\text{Ce}^{3+}]_t = [\text{Ce}^{3+}]_0 \exp(-kt)$ expressed as $(100 - A)\exp(-k_1t) + A$ on a 100 basis to take into account the A nonreacting cerium fraction. A is the threshold of the minimum intensity reached over a long time, and k_1 is the constant velocity (s^{-1}) of cerium III photooxidation. The results are shaped as green curves in Fig. 7b and c. (ii model) The reaction is considered a second-order mechanism, always relying on a single redox centre. V is calculated as follows: $-d[\text{Ce}^{3+}]/dt = k_2[\text{Ce}^{3+}]^2$, equal to $(100 - A)/(1 + 2(100 - A)) \times k_2t + A$, where k_2 is the two-order velocity constant (s^{-1}) (blue curves). Finally, the accumulation of two simultaneous 1st order kinetic laws (iii model) (pink curves) is such as $-d[\text{Ce}^{3+}]/dt = k_1[\text{Ce}^{3+}]^1 + k_2[\text{Ce}^{3+}]^2$ leading to a disappearance kinetic such as $(100 - A - B)\exp(-k_1t) + B\exp(-k_2t) + A$, with k_1 and k_2 used for the slow and fast kinetic constants, respectively. The two first law models are not fully satisfactory, and significant residual signals are obtained at the beginning of the irradiation and at 16 hours and 4 hours for Rb_2K and K_3 , respectively. However, the (iii model) leads to a very low R^2 value, and it is reasonable to conclude that two different Ce^{3+} sites with two different kinetics are involved. The total fitted parameter values are reported in Table S2 (ESI[†]).

The most surprising finding is that the main part of the cerium is actively embedded in the redox process since the A nonreacting fraction is approximately 20–30% (28.13 and 18.34 are the values extracted from the fit on Rb_2K and K_3 , respectively). Indeed, we have shown in our previous work in Rb_2K compound that the redox is clearly originating from an overlapping of the cerium and indium external atomic orbitals,

i.e., from cations in the first neighbouring positions. The redox process involves two Ce^{3+} ions for one In^+ cation, as illustrated below:



The probability of being involved in a redox process for Ce^{3+} ions, *i.e.*, to not be “isolated” from other Ce^{3+} ions is $P = 1 - (1 - p)^n$, where p is the substitution rate by Ce^{3+} at the indium site, and n is the total amount of potential locations to find cerium cations that can coparticipate in the redox reaction (active redox coordination sphere number). The n value is equal to 56 because, to be isolated from redox potentiality, each Ce^{3+} ion must be placed farther from a next-to-next neighbouring position from a second Ce^{3+} ion (56 next-to-next neighbours).²⁰ The probability of finding a second active cerium ion around the one considered in the active sphere is $1 - 0.98^{56} \approx 68\%$ for $p = 2\%$, in relatively good agreement with the A value (nonreactive cerium fraction). This first calculation indicates that the redox mechanism is effective, while two cerium cations belong to the first neighbouring sphere of a single indium site (with 12 neighbours). Furthermore, electron transfer between two cerium III must also be considered helpful for this process. Additionally, it can be imagined that Ce^{3+} – Ce^{3+} – In^{3+} chains within a “cascade” double electron injection from the two cerium ions towards the indium site, occur more quickly than the case that two independent cerium ions have to inject one electron each into a “central” indium site. Then, B eliminates the cerium rate, which is the proportion of ions concerned by the rapid kinetic law (k_2 constant) and can correspond to cerium ions involved in this “cascade” transfer. Considering a doping with exactly 2% of cerium on indium site with perfectly homogeneous distribution, the iii model, implemented with consideration of a quick (Ce–Ce–In triplets) and slow (Ce–In–Ce triplets) redox centre mechanisms, proportion of nonreacting cerium ions (A fraction)/reacting with fast kinetic (B fraction)/

reacting with slow kinetic ($100 - A - B$ fraction) will be: $1 - (1 - 0.98^{56}) = 32\%/1 - (0.98^{12} \times (1 - 0.98^{56})) = 46.6\%/100 - 46.6 - 32 = 21.4\%$, respectively. These results are in good agreement with the experimental values.

After far UV irradiation, the luminescence of monovalent indium cations was detected for the three studied compounds. It consists of $5s^2 \leftrightarrow 5s^15p^1$ transitions. Concerning the In^+ cation, spin–orbit coupling results in the global splitting of the p orbitals into 3P_0 , 3P_1 , 3P_2 and 1P_1 levels, with 1S_0 being the ground state. The absorption bands are labelled A (1S_0 – $^3P_{0,1}$), B (1S_0 – 3P_2), and C (1S_0 – 1P_1).²¹ In an octahedral environment, the emission results from the de-excitation of the 3P_1 level, which is split into a $^3A_{1u}$ (A_T high energy component) and a $^3A_{1u}$ (A_x low energy component). The first transition is usually quenched at room temperature. In addition, the Jahn–Teller effect is predominant among the spin–orbit couplings, and the spectral distribution in absorption and emission is strongly affected Fig. 8 shows the excitation and emission spectra of the three fluoride compounds.^{22–25}

As indicated above, the redox process induced under UV irradiation stabilized indium in the monovalent oxidation state. In the three fluorides, a large orange-red emission due to the A_x (3P_0) \rightarrow 1A_1 (1S_0) transition is observed, and the corresponding excitation bands peak at approximately 240 nm. For the Rb_2K and K_3 compounds, measurements were performed at 8 K to confirm the origin of these transitions through the expected appearance of the A_T component. The spectral characteristics are listed in Table 4.

Both excitation and emission bands are redshifted in parallel to the decrease in the alkali radii. This reflects the Nephelauxetic effect first described by C. K. Jørgensen in 1962.²⁶ The interelectronic repulsion of a transition metal is weaker when integrated in a complex than for the free ion because of the overlap of the ligand–metal orbitals and the expansion of the electronic cloud over a wider area. The electronic repulsion is reduced, leading to decreased energy levels.^{27,28} This effect is driven by the covalence and polarizability of the chemical bonding. The excitation energy decreases with increasing bond covalence.

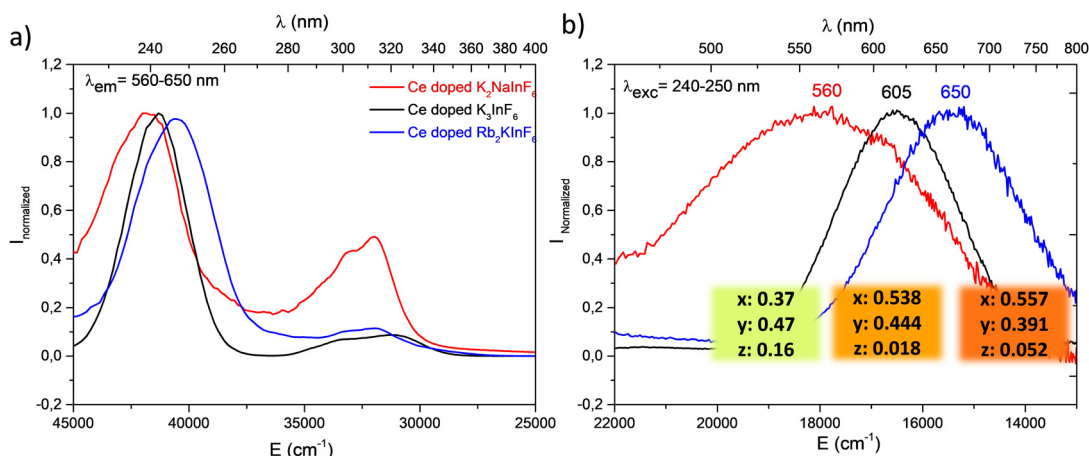
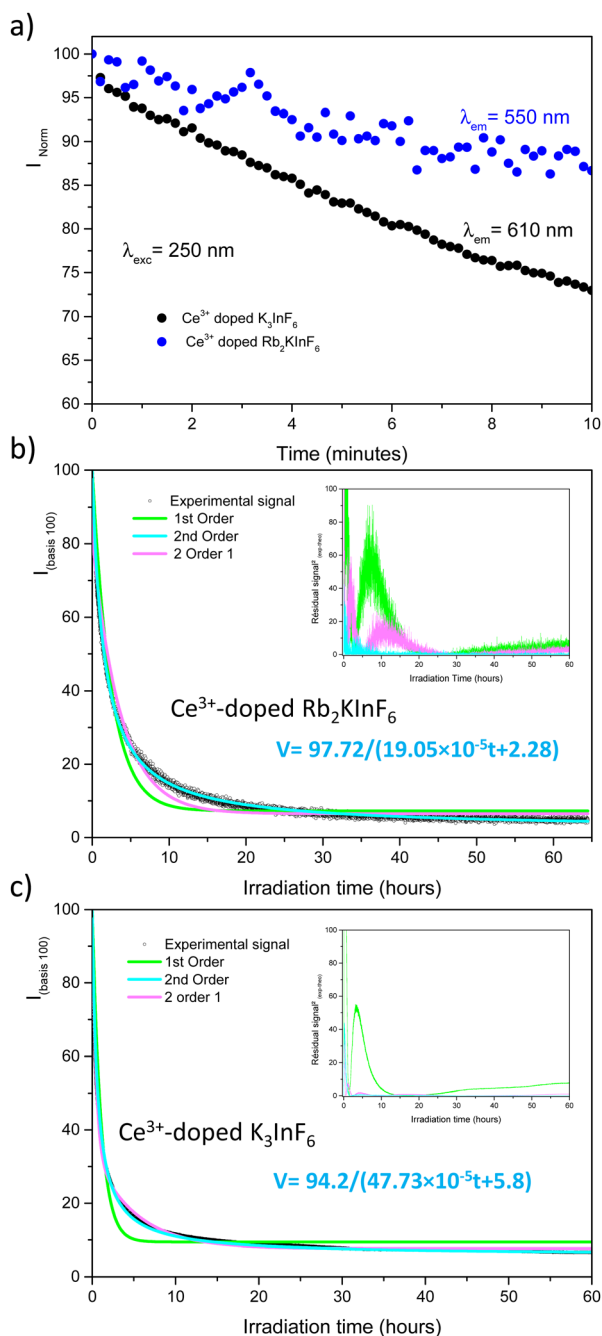


Fig. 8 Excitation (a) and emission spectra (b) after 10 minutes of irradiation at 315 nm of Ce^{3+} doped fluoride compounds. Measurements were performed at room temperature.



	λ_{ex} max (cm ⁻¹)	λ_{em} max (cm ⁻¹)			ΔE (cm ⁻¹)	
Compound	$T = 293$ K	A_T (293 K)	A_X (293 K)	A_T (8 K)	A_X (8 K)	$\Delta(^3\text{P}_{0,1}(A_X)-^1\text{S}_0)$ (cm ⁻¹) 293 K $\Delta(A_T - A_X)$ (cm ⁻¹) 8 K
Rb ₂ K [#] ref. 8	40 485 (247 nm)	—	15 385 (650 nm)	25 000 [#] (400 nm)	15 152 [#] (660 nm)	25 100
K ₂ Na	42 017 (238 nm)	—	17 587 (560 nm)	—	—	24 430
*K ₃	41 322 (242 nm)	380	16 526 (605 nm)	26 316 (380 nm)	15 873 (630 nm)	24 800
						10 443



Finally, the UV irradiation of Ce^{3+} -doped Rb_2KInF_6 , K_2NaInF_6 elpasolites and K_3InF_6 cryolite makes possible a reversible switch of the luminescence from blue to yellow-orange-red from a reversible redox process between cerium

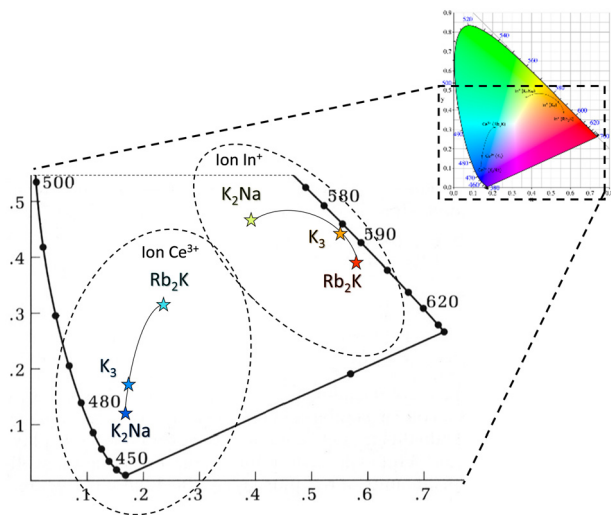


Fig. 10 CIE xy trichromatic diagram based on Ce^{3+} and In^{+} in Ce^{3+} -doped Rb_2KInF_6 , K_2NaInF_6 elpasolites and K_3InF_6 cryolite.

and indium ions due to atomic orbital overlapping. Due to different polarizabilities, the use of various alkali cations makes it possible to tune the redox kinetics and these emissive colours, as illustrated in Fig. 10.

From the chromatic x and y parameters between the Ce^{3+} and In^{+} spectral distributions, the maximum of the calculated optical contrast ($\text{OC} = \sqrt{(x_{\text{Ce}^{3+}} - x_{\text{In}^{+}})^2 + (y_{\text{Ce}^{3+}} - y_{\text{In}^{+}})^2}$) is obtained with the cryolite compound and is equal to 0.482. OC values of 0.314 and 0.424 are calculated for the Rb_2K and K_2Na compounds, respectively.

4. Conclusion

From solid-state synthesis, three Ce^{3+} -doped fluorides, Ce -doped Rb_2KInF_6 and K_2NaInF_6 elpasolite-type and Ce -doped K_3InF_6 cryolite-type, have been prepared. In addition to the refinement of the crystallographic elpasolite structures as already well described in the literature, coupling X-ray diffraction and Raman spectroscopy, a new space group with full atomic positioning was herein proposed for the cryolite-type compound.

The reversible photochromoluminescent properties of Ce^{3+} -doped Rb_2KInF_6 , K_2NaInF_6 elpasolite and Ce -doped K_3InF_6 cryolite have been demonstrated. Supported by the neighbouring probability of the involved cations, photoluminescence is confirmed to come from the redox process between neighbouring cerium and indium ions due to orbital overlap. Before far UV irradiation ($\text{Ce}^{3+}/\text{In}^{3+}$ state), Ce^{3+} emission bands are shifted to lower energy with increasing alkali size. After the forth-redox reaction ($\text{Ce}^{4+}/\text{In}^{+}$ state), the In^{+} luminescence is also tuned by the alkali size due to the Nephelauxetic effect. Hence, cyan to red (Rb_2K) or blue to yellow (K_3) photochromism can be achieved. A deep kinetic investigation was performed to better understand the successive decrease in Ce^{3+} and In^{+} emission during forth- and back-irradiation, respectively. The forth-redox is governed by

the accumulation of two 1st order laws, while for the back-redox, the In^{+} disappearance kinetic curve follows a 2nd order law. The redox processes appear faster for the cryolite compound than for the Rb_2K compound, suggesting that a more rigid host lattice accelerates electron transfer. Noticeable changes in the optical contrasts were obtained along the fluoride series.

Conflicts of interest

There are no conflicts to declare.

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References

- Y. Hirshberg, Reversible Formation and Eradication of Colors by Irradiation at Low Temperatures. A Photochemical Memory Model, *J. Am. Chem. Soc.*, 1956, **78**(10), 2304, DOI: [10.1021/ja01591a075](https://doi.org/10.1021/ja01591a075).
- G. P. Smith, Photochromic glasses: Properties and applications, *J. Mater. Sci.*, 1967, **2**, 139, DOI: [10.1007/BF00549573](https://doi.org/10.1007/BF00549573).
- R. Zhang, *The Syntheses, NMR and Photochromic Properties of Modified Dimethyldihydropyrenes*, PhD thesis, University of Victoria 2002, <https://hdl.handle.net/1828/247>.
- H. J. Hoffmann, The use of silver salts for photochromic glasses, *Stud. Org. Chem.*, 2003, **40**, 822.
- H. Rosset, *Security element having a variable optical effect and security sheet or document or article comprising it*, FR2933428, 2010.
- W. J. Cross and W. J. Hillebrand, *Beitr. Krystallogr. Mineral.*, 1887, **12**, 495.
- J. P. Chaminade, A. Garcia, T. Gaewdang, M. Pouchard, J. Grannec and B. Jacquier, Reversible photoionization process in luminescent Ce^{3+} doped elpasolite-type fluoroindates, *Radiat. Eff. Defects Solids*, 1995, **135**(1-4), 137, DOI: [10.1080/10420159508229823](https://doi.org/10.1080/10420159508229823).
- L. Cornu, M. Gaudon, O. Toulemonde and P. Veber, Optical contrast and cycling of bistable luminescence properties in $\text{Rb}_2\text{KIn}(1-x)\text{Ce}x\text{F}_6$ compounds, *Dalton Trans.*, 2016, **45**, 3380, DOI: [10.1039/C5DT04772B](https://doi.org/10.1039/C5DT04772B).
- L. Cornu, M. Gaudon, P. Veber, A. Villesuzanne, S. Pechev, A. Garcia and V. Jubera, Discussion on the Structure Stability and the Luminescence Switch under Irradiation of a Ce-Doped Elpasolite Compound, *Chem. – Eur. J.*, 2015, **21**(13), 5242, DOI: [10.1002/chem.201405784](https://doi.org/10.1002/chem.201405784).
- G. Blasse and B. C. Grabmaier, *Luminescent materials*, Springer-Verlag; Berlin Heidelberg New York, 1994, London



- Paris Tokyo, Hong Kong Barcelona Budapest, ISBN: 978-3-642-79017-1, DOI: [10.1007/978-3-642-79017-1](https://doi.org/10.1007/978-3-642-79017-1).
- 11 Y. Mao, H. Zhou and S. S. Wong, Synthesis, Properties, and Applications of Perovskite-Phase Metal Oxide Nanostructures, *Mater. Matters*, 2010, **50**, 1.
 - 12 H.-R. Wenk and A. G. Bulach, *Minerals: their constitution and origin*, Cambridge Univ. Press, 2004.
 - 13 D. Babel, R. Haegle, G. Pausewang and F. Wall, Ueber kubische und hexagonale elpasolithe $\text{Al}_2\text{BIMIIIIF}_6$, *Mater. Res. Bull.*, 1973, **8**(12), 1371, DOI: [10.1016/0025-5408\(73\)90021-4](https://doi.org/10.1016/0025-5408(73)90021-4).
 - 14 H. Bode and E. Voss, Strukturen der Hexafluorometallate(III), *Z. Anorg. Allg. Chem.*, 1957, **290**(1-2), 1, DOI: [10.1002/zaac.19572900102](https://doi.org/10.1002/zaac.19572900102).
 - 15 K. S. Aleksandrov, S. V. Misyul, M. S. Molokeev and V. N. Voronov, Structures of distorted phases and critical and noncritical atomic displacements of elpasolite Rb_2KInF_6 during phase transitions, *Phys. Solid State*, 2009, **51**, 2505.
 - 16 K. Nakamoto, *Infrared and Raman Spectra of Inorganic and Coordination Compounds: Part B: Applications in Coordination, Organometallic, and Bioinorganic Chemistry*, 2008, DOI: [10.1002/978047](https://doi.org/10.1002/978047).
 - 17 W. M. Yen, S. Shionoya and H. Yamamoto, *Phosphor handbook*, Boca Raton, FL: CRC Press/Taylor and Francis, 2nd edn, 2007.
 - 18 https://fr.wikiversity.org/wiki/Cin%C3%A9tique_chimique/Lois_de_vitesse_simples.
 - 19 <https://www.techno-science.net/definition/3325.html>.
 - 20 L. Cornu, Matériaux X-Chromo-Luminescent de type Spinelle et elpasolite: relation structure-propriétés, thèse de doctorat en chimie, *Univ. Bordeaux*, 2015, 49–179.
 - 21 A. Fukuda, Jahn-Teller Effect on the Structure of the Emission Produced by Excitation in the A Band of KI: Tl-Type Phosphors. Two Kinds of Minima on the $\Gamma_4-(3T_{1u})$ Adiabatic Potential-Energy Surface, *Phys. Rev. B: Solid State*, 1970, **1**(10), 4161, DOI: [10.1103/PhysRevB.1.4161](https://doi.org/10.1103/PhysRevB.1.4161).
 - 22 D. Klick and H. G. Drickamer, High-pressure studies of Jahn-Teller-split luminescence in alkali halides doped with In^+ and Tl^+ , *Phys. Rev. B: Solid State*, 1978, **17**(3), 952, DOI: [10.1103/PhysRevB.17.952](https://doi.org/10.1103/PhysRevB.17.952).
 - 23 A. Fukuda, Magnetic Field Effect on the Triplet Relaxed Excited States Responsible for the AT and AX Emission bands of Ga^+ and In^+ Centers in Alkali Halides, *J. Phys. Soc. Jpn.*, 1976, **40**(3), 776, DOI: [10.1143/JPSJ.40.776](https://doi.org/10.1143/JPSJ.40.776).
 - 24 A. I. Popov and I. Plavina, Photostimulated emission of KBr-In previously exposed to UV- or X-radiation, *Nucl. Instrum. Methods Phys. Res., Sect. B*, 1995, **101**, 252.
 - 25 V. S. Sivasankar, K. Schmitt and P. W. M. Jacobs, Luminescence and decay times of CsI:In^+ , *J. Lumin.*, 1985, **33**(4), 409, DOI: [10.1016/0022-2313\(85\)90110-3](https://doi.org/10.1016/0022-2313(85)90110-3).
 - 26 C. K. Jørgensen, Absorption spectra of transition group complexes of sulphur-containing ligands, *J. Inorg. Nucl. Chem.*, 1962, **24**(12), 1571, DOI: [10.1016/0022-1902\(62\)80011-6](https://doi.org/10.1016/0022-1902(62)80011-6).
 - 27 C. K. Jørgensen, Spectroscopy of Transition-Group Complexes, in *Advances in Chemical Physics*, ed. I. Prigogine, John Wiley & Sons, Inc., Hoboken, NJ, USA, 2007, p. 33.
 - 28 A. L. Tchougréeff and R. Dronskowski, Nephelauxetic effect revisited, *Int. J. Quantum Chem.*, 2009, **109**(11), 2606, DOI: [10.1002/qua.21989](https://doi.org/10.1002/qua.21989).

