Photoluminescence of RbCaF₃:Mn²⁺: the influence of phase transitions

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Abstract. Precise photoluminescence measurements on an RbCaF₃:Mn²⁺ sample containing only 400 ppm of Mn²⁺ have been carried out in the 10-300 K temperature range. The results are compared with those obtained in other fluoroperovskites doped with Mn²⁺. The analysis of the ${}^6A_{1g}(S) \rightarrow {}^4T_{1g}(G)$ excitation peak at room temperature leads to a Mn²⁺-F⁻ distance R=213.3 pm which is close to that derived from the experimental isotropic superhyperfine constant, A_s . The plot of the first moment of the emission band, M_1 , against temperature reveals a slight but sensible change of slope at T=193 K which is associated with the $O_b^1 \rightarrow D_{4b}^{18}$ structural phase transition of the host lattice. Furthermore, at T=40 K, M_1 undergoes an abrupt increase of ~ 100 cm⁻¹. This fact supports the existence of mother phase transition involving an increase $\Delta R/R \simeq 0.2\%$ upon cooling, and thus a situation which is similar to that detected in the structural phase transition of KMnF₃ at $T_{c3}=81.5$ K. To our knowledge this is the first time that clear evidence of both phase transitions in RbCaF₃ has been achieved through an optical probe. Finally the variation of the ${}^4A_{1g}(G)$, ${}^4E_g(G)$ peak, E_3 , along the fluoroperovskite series is analysed.

1. Introduction

Some cubic fluorides doped with Mn^{2+} are among the few systems where a local structural characterization around the impurity has properly been achieved. As an example the Mn^{2+} - F^- distance, R, between the substitutional Mn^{2+} impurity and the nearest F^- ions has been determined for $KZnF_3:Mn^{2+}$ and $RbCdF_3:Mn^{2+}$ using three independent methods: (1) the extended x-ray absorption fine structure (EXAFS) technique [1]; (2) the analysis of the isotropic superhyperfine (SHF) constant, A_s [2, 3]; and (3) the analysis of excitation spectra obtained through photoluminescence [4, 5]. In this line very recent EXAFS measurements on $CaF_2:Mn^{2+}$ lead to R=227 pm [6] which is coincident with previous determinations from the experimental A_s value [3].

As a common feature for isolated Mn^{2+} in cubic fluorides it has been demonstrated that crystal-field spectra and electron paramagnetic resonance (EPR) parameters can reasonably be understood only on the basis of isolated MnF_N units whose equilibrium distance, R, is influenced by the chemical pressure exerted by the rest of the lattice. In the series of cubic fluoroperovskites (at room temperature) such a pressure orders the R values as the corresponding values, R_0 , for the perfect lattice although the two figures are usually different for a given lattice. This situation is reviewed in [7].

Although a great many cubic fluoroperovskites doped with Mn^{2+} have been investigated through EPR [8] (and some of them also by electron-nuclear double resonance (ENDOR) [9–11]) the number of such systems explored using photoluminescence techniques is certainly smaller. This is partly because of the very small oscillator strengths ($f \simeq 10^{-8}$) associated with crystal-field transitions of isolated MnF_6^{4-} units in cubic symmetry which makes it difficult to detect photoluminescence spectra when the Mn^{2+} concentration is below about 1000 ppm.

The present work is devoted to exploration of the luminescence and excitation spectra of an RbCaF₃:Mn²⁺ sample containing only about 400 ppm of Mn²⁺ as well as the associated lifetime in the 10 K-300 K range. Apart from detecting the photoluminescence due to MnF₆⁴ a first goal of our investigation is to explore whether or not the crystal-field spectrum of RbCaF₃:Mn²⁺ is compatible with an Mn^{2+} -F⁻ distance R = 214.2(6) pm as previously derived from the analysis of the experimental A_s value [2]. Other goals of the present work are related to phase transitions in RbCaF₃. This compound has a clearly established $O_h^1 \rightarrow D_{4h}^{18}$ structural phase transition at $T_{cl} = 193 \text{ K} [12-18]$. This phase transition is also observed in RbCdF₃ ($T_c = 124$ K) [19] but not in CsCaF₃ [20]. EPR results on RbCaF₃:Mn²⁺ and RbCdF₃:Mn²⁺ [8, 21], and specially ENDOR data on the latter system [11] indicate that the local geometry around Mn²⁺ is practically octahedral in the tetragonal phase. To be more specific, if $R_{\rm ax}$ and $R_{\rm eq}$ denote respectively the axial and equatorial Mn²⁺-F distances of the distorted MnF₆⁴ unit in the tetragonal phase, it has been found that $R_{\rm ax}-R_{\rm eq}=0.2$ pm for RbCdF₃:Mn²⁺ at T=30 K [7]. This value has to be compared with $R_{\rm ax}^0 - R_{\rm eq}^0 = 1.1$ pm corresponding to the perfect lattice [19]. A similar situation would occur for RbCaF3:Mn2+ as discussed in [7]. Thus if changes in the optical spectrum of Mn²⁺ in fluorides mainly reflect geometrical changes in the first coordination sphere the detection of the phase transition of RbCaF₃ at $T_{\rm cl} = 193$ K using optical parameters of the Mn²⁺ impurity as probe is not in principle a simple task. Recent results by Villacampa et al on RbCdF₃:Cr³⁺ [22] supported this view. In order to demonstrate the influence of this phase transition upon optical parameters associated with MnF₆⁴⁻ in RbCaF₃:Mn²⁺, measurements on CsCaF₃:Mn²⁺ under the same experimental conditions have also been performed.

As a last goal this work is also devoted to explore the existence of another phase transition in RbCaF₃ at $T_{c2} \cong 40$ K. Evidence of such a phase transition has been obtained through early heat capacity, dielectric and Raman studies [12, 13] as well as by subsequent x-ray and dielectric measurements by Hidaka et al [18]. Nevertheless no indication of this phase transition has been achieved through the neutron measurements carried out by Kamitakahara and Rotter [14]. Also other studies devoted to exploring the first phase transition at $T_{c1} = 193$ K seem to have paid no attention to the second one [15–17, 21] and so it thus deserves further investigation.

2. Experimental details

The single crystals of RbCaF₃: Mn^{2+} used in this work were kindly provided by Professor R Alcalá. The actual Mn^{2+} concentration, 400 ± 100 ppm, was estimated by measuring the EPR integrated intensity at RT of a powder (obtained from the sample) and comparing it to the corresponding value for a $KZnF_3$: Mn^{2+} powder with 2% Mn^{2+} . Qualitatively speaking the low concentration of Mn^{2+} in our RbCaF₃ sample

is also supported by the well resolved SHF structure seen in its EPR spectrum. That structure is however fully absent for the KZnF₃:Mn²⁺ powder where the average distance between closest Mn²⁺ would be about 3.7 times smaller than in our sample. The value of the isotropic SHF constant $A_s = 14.4(4) \times 10^{-4}$ cm⁻¹ measured at RT for RbCaF₃:Mn²⁺ is essentially coincident with $A_s = 13.9(3) \times 10^{-4}$ cm⁻¹ previously reported by Rousseau *et al* [8].

For obtaining excitation and emission spectra of RbCaF₃:Mn²⁺ as well as for carrying out lifetime measurements it was necessary to improve the detection sensitivity because of the low value of the Mn²⁺ concentration.

Excitation spectra were obtained with an implemented Jobin Yvon JY-3D fluorimeter [23] equipped with a photon counting facility. The excitation beam was chopped and focused on the sample which was coupled to an Oriel interference filter ($\lambda = 600$ nm) in front of a Hamamatsu R928 photomultiplier. The luminescence signal was measured during the non-excitation periods with a Stanford System SR440 preamp and SR 400 photon counter.

Luminescence spectra were obtained by means of a John Yvon HR-320 monochromator employing the chopping light of a Spectra Physics model 2020-03 Ar laser as excitation source.

For lifetime measurements, the luminescence signal was digitized by means of a Tektronix 2430A scope.

Temperature variations were achieved by a Scientific Instruments DE-202 closed-circuit cryostat allowing temperature stabilities within 0.1 K and an accuracy of 0.5 K, with an APD-K temperature controller.

3. Results and discussion

The RT luminescence of RbCaF₃:Mn²⁺ together with the corresponding excitation spectrum are shown in figure 1 where spectra of CsCaF₃:Mn²⁺ are also included for comparison. As in other fluorides doped with Mn²⁺ the only emission band found in RbCaF₃:Mn²⁺ is assigned as the ${}^4T_{1g}(G) \rightarrow {}^6A_{1g}(S)$ electronic transition thus involving the *first* excited state. The assignment of excitation bands for RbCaF₃:Mn²⁺ together with the position of the corresponding maxima are given in table 1. Values of the effective Racah parameters, B and C, together with that for the cubic field splitting parameter, 10Dq, are also given.

The excitation spectrum of RbCaF₃:Mn²⁺ is very similar but not identical to that of CsCaF₃:Mn²⁺ as can be inferred simply by looking at figure 1. The most significant difference between both spectra lies in the position of the first $^6A_{lg}(S) \rightarrow {}^4T_{lg}(G)$ excitation peak which is found at $\lambda = 511$ nm for RbCaF₃:Mn²⁺ while it appears at $\lambda = 503$ nm for CsCaF₃:Mn²⁺. This result concurs with previous work on Mn²⁺ doped fluorides showing that among the observed crystal-field transitions of MnF₆⁴⁻ the $^6A_{lg}(S) \rightarrow {}^4T_{lg}(G)$ is the most sensitive one to a change of the host lattice [4, 5]. Microscopically this relevant fact has been explained on the following grounds:

- (i) The near independence of the effective Racah parameters B and C from R observed experimentally and well supported by theoretical calculations on MnF_6^{4-} in fluoroperovskites performed for several values of R [24].
- (ii) The ${}^6A_{lg}(S) \rightarrow {}^4T_{lg}(G)$ transition is the most strongly dependent upon 10Dq among the observed crystal-field transitions [25].

Table 1. Experimental peak energy and assignment of the crystal field bands observed in the room temperature excitation spectra of RbCaF₃:Mn²⁺ and CsCaF₃:Mn²⁺, together with the fitted values of the Racah parameters B and C, and the crystal field 10Dq. Tree's ($\alpha = -65$ cm⁻¹) and seniority ($\beta = 131$ cm⁻¹) corrections were included in the fitting. σ is the standard deviation. The position of the luminescence band is also included. Units are in cm⁻¹. The errors in the peak energies and 10Dq values are 20 cm⁻¹ and 40 cm⁻¹, respectively.

Assignment	RbCaF ₃ :Mn ²⁺	CsCaF ₃ :Mn ²⁺ 19 880	
$\overline{{}^{4}T_{1g}(G) \leftarrow {}^{6}A_{1g}(S)}$	19 560		
⁴ T _{2g} (G)	23 650	23 805	
${}^{4}A_{1g}(G), {}^{4}E_{g}(G)$	25 400	25 350	
$^{4}T_{2g}(D)$	28 450	28 500	
$^{4}E_{g}(D)$	30 280	30 265	
$^{4}T_{1g}(P)$	32 340	31 960	
\boldsymbol{B}	831	834	
C	3158	3141	
10Dq	7305	6890	
σ	215	225	
Emission: ${}^{4}T_{1g}(G) \rightarrow {}^{6}A_{1g}(S)$	17 920	18 350	

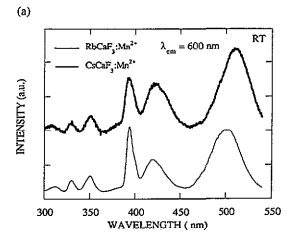
(iii) The significant dependence of 10Dq upon R found experimentally [4,5,26] as well as by means of theoretical calculations [24,27] on transition-metal complexes as a function of R. This dependence can be written as

$$10Dq = KR^{-n}. (1)$$

Experimentally [4, 5] a value n=4.7 was found for MnF_6^{4-} . The validity of these ideas is well supported by the general view on the main optical results for Mn^{2+} doped fluoroperovskites given in table 2. Judging by these the influence of the so-called explicit temperature contribution [28, 29] to the band maxima appears to be not important as regards the comparison among all Mn^{2+} doped fluoroperovskites at RT. By contrast it plays a relevant role for understanding the temperature dependence of the peak energies for a given system [29].

Table 2. Metal-ligand distance and spectroscopic parameters in the Mn^{2+} doped ABF₃ fluoroperovskite series at room temperature. R_0 is the B^{2+} - F^- distance in the host lattice. $R_{\rm C}$ is the Mn^{2+} - F^- distance determined by EXAFs in KZnF₃: Mn^{2+} and RbCdF₃: Mn^{2+} , and through the isotropic superhyperfine constant, $A_{\rm s}$, in KMgF₃: Mn^{2+} , RbCaF₃: Mn^{2+} , and CsCaF₃: Mn^{2+} . R^4 - $R^$

System	R_0	⁴ T _{1g} (G)	⁴ A _{1g} , ⁴ E _g (G)	Emission	10 <i>Dq</i>	R _C [7]	R4 _{T12}	R_{10Dq}
KMgF ₃ :Mn ²⁺	199.3	18 160	25 200	16 955	8432	207.0	206.0	206.8
KZnF ₃ :Mn ²⁺	202.6	18 530 [37]	25 210 [37]	17 100 [37]	8215	208.0	208.0	208.0
KMnF ₃	209.5	18 900 [29]	25 245 [29]	_ ` ´	7853	209.5	209.9	209.9
RbMnF ₃	212.0	19 300 [29]	25 275 [29]	_	7496	212.0	211.9	212.0
RbCdF ₃ :Mn ²⁺	220.0	19 510	25 215	17 950	7264	213.0	213.0	213.4
RbCaF ₃ :Mn ²⁺	222.7	19 560	25 400	17 920	7306	214.2	213.3	213.1
CsCaF ₃ :Mn ²⁺	226.2	19 880	25 350	18 350	6890	215.5	214.9	215.8



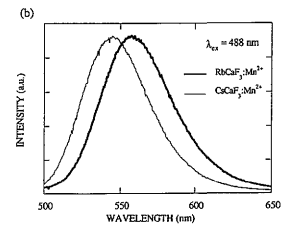


Figure 1. Room temperature (a) excitation and (b) luminescence spectra corresponding to RbCaF₃: Mn^{2+} and CsCaF₃: Mn^{2+} . A small shoulder observed in the second excitation band peaked at ~ 420 nm comes from xenon lines of the lamp.

As regards RbCaF₃:Mn²⁺ itself within the series of fluoroperovskites doped with Mn²⁺ the position of the $^6A_{lg}(S) \rightarrow ^4T_{lg}(G)$ excitation peak at RT lies between that for RbCdF₃:Mn²⁺ and that for CsCaF₃:Mn²⁺ as is expected provided R values along the series are *ordered* as the corresponding R_0 values. However, the small difference between the excitation peaks of RbCdF₃:Mn²⁺ and RbCaF₃:Mn²⁺ indicates that the corresponding R values are close to each other. This idea is also supported by the analysis of the corresponding 10Dq values which give rise however to an R value slightly higher for RbCdF₃:Mn²⁺ than for RbCaF₃:Mn²⁺.

For achieving a proper comparison between the R values of these systems we have also performed EPR measurements on a powder of RbCdF₃:Mn²⁺ where the SHF structure is clearly visible. The measured value $A_s = 15.7(3) \times 10^{-4}$ cm⁻¹ is higher than $A_s = 14.4(4) \times 10^{-4}$ cm⁻¹ measured for RbCaF₃:Mn²⁺. Such figures imply [2] that the difference, ΔR , between R values corresponding to RbCaF₃:Mn²⁺ and RbCdF₃:Mn²⁺ would be $\Delta R = (1.8 \pm 1.2)$ pm, and so R values would be

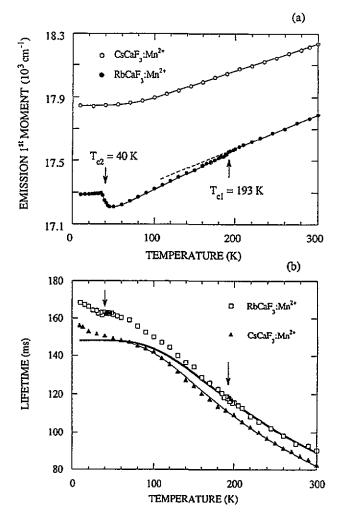


Figure 2. Plot of (a) the experimental luminescence first moment, M_1 , and (b) the lifetime τ against temperature for RbCaF₃:Mn²⁺ and CsCaF₃:Mn²⁺. In the case of the lifetime the solid lines have been obtained through a fitting of the experimental values using equation (3). The fitted parameter values are $\hbar\omega_u=260~{\rm cm}^{-1},~\tau_0=148~{\rm ms}$ for CsCaF₃:Mn²⁺ and $\hbar\omega_u=290~{\rm cm}^{-1},~\tau_0=148~{\rm ms}$ for RbCaF₃:Mn²⁺. Lines in $M_1(T)$ are included as visual guides.

ordered as the corresponding R_0 values as inferred from the experimental A_s .

The main reason why the fitted 10Dq leads to a slightly higher R value for RbCdF₃:Mn²⁺ than for RbCaF₃:Mn²⁺ comes from the complex structure of the asymmetric ${}^6A_{1g}(S) \rightarrow {}^4A_{1g}(G)$, ${}^4E_g(G)$ band. In fact 10Dq is essentially determined by the difference $\Delta = E_3 - E_1$ between the energies of the third $({}^6A_{1g}(S) \rightarrow {}^4A_{1g}(G), {}^4E_g(G))$ and first $({}^6A_{1g}(S) \rightarrow {}^4T_{1g}(G))$ maxima. Nevertheless, problems appear in the interpretation of the third band maximum because (i) the two ${}^6A_{1g}(S) \rightarrow {}^4A_{1g}(G)$ and ${}^6A_{1g}(S) \rightarrow {}^4E_g(G)$ transitions are neither exactly coincident (they are separated by 100 cm^{-1} in both KMgF₃:Mn²⁺ and KZnF₃:Mn²⁺ [30]) nor display the same

oscillator strength, and (ii) the associated vibronic structure can change for different systems.

In general, the vibronic couplings responsible for both the electric dipole oscillator strength and the phonon sideband of the crystal field transitions may be influenced not only by changes in R but mainly by the type of cations out of the first coordination sphere. In this way, both the phonon frequency and the electronphonon coupling associated with the vibronic sidebands of the ${}^6A_{1g}(S) \to {}^4A_{1g}(G)$ and ${}^6A_{1g}(S) \to {}^4E_g(G)$ transitions as well as of the ${}^6A_{1g}(S) \to {}^4T_{1g}(G)$ one are indeed rather different in KMgF3:Mn2+ and KZnF3:Mn2+. Experimental evidence of these facts can be found elsewhere [30, 31]. The $T=1.7~{\rm K}$ high-resolution spectra around the ⁴A_{1g}, ⁴E_g(G) in KMgF₃:Mn²⁺ and KZnF₃:Mn²⁺ show that the dipolar magnetic zero-phonon lines are shifted 100 cm⁻¹ towards higher energies in KMgF₃:Mn²⁺ [30] at variance with the RT situation in which the band maximum for KMgF₃:Mn²⁺ lies only 10 cm⁻¹ below the KZnF₃:Mn²⁺ one [5]. This result, which cannot be explained on the basis of different values of R, clearly suggests that vibronic couplings play a crucial role in determining the bandshape and therefore the band maxima in such non-10Dq-dependent ${}^4A_{1g}$, ${}^4E_e(G)$ tansitions. Such an effect could also justify the anomalous dispersion of the ${}^4A_{1g}$, ${}^4E_g(G)$ maxima, E_3 , observed along the fluoroperovskite series whose variation, as shown in table 2, cannot be understood simply by changes in the Mn²⁺-F⁻ distance. Taking into account the measured value $\partial E_3/\partial R = -15$ cm⁻¹ pm⁻¹ for RbMnF₃ [32], E_3 should increase by $\sim 120 \text{ cm}^{-1}$ if R increases by $\sim 8 \text{ pm}$. Though this increase reflects the trend followed by E_3 , this mechanism is unable to explain its variation between successive crystals along the series. This is particularly puzzling on going from RbCdF3:Mn2+ to RbCaF₃:Mn²⁺ and CsCaF₃:Mn²⁺ where blue shifts of 185 and 135 cm⁻¹ are actually observed instead of the 30 and 50 cm⁻¹ that should be expected from their respective R values.

This analysis suggests that spectroscopic studies of the R dependence of E_3 should be performed on a given crystal using hydrostatic pressure techniques and not chemical pressures where the use of different host matrices affects not only R but also the vibronic couplings. This effect is however not so important for the 10Dq dependent $^6A_{1g}(S) \rightarrow {}^4T_{1g}(G)$ excitation band, E_1 , whose R dependence, $\partial E_1/\partial R = 200$ cm⁻¹ pm⁻¹, is one order of magnitude higher than that of E_3 .

Owing to this, we have fitted the experimental position of ${}^6A_{1g}(S) \rightarrow {}^4T_{1g}(G)$ maxima corresponding to fluoroperovskites containing Mn²⁺ to the equation:

$$E_1 = a + bR. (2)$$

The constants a and b have been determined using the values of E_1 and R corresponding to KMnF₃ and RbMnF₃ as well as those for KZnF₃:Mn²⁺ and RbCdF₃:Mn²⁺. In the latter cases the distance, R, was determined through EXAFS [1]. The best values of the constants a and b were found to be a=21875 cm⁻¹ and b=194.3 cm⁻¹ pm⁻¹. The actual Mn²⁺-F⁻ distance obtained for the other fluoroperovskites (CsCaF₃, RbCaF₃ and KMgF₃) doped with Mn²⁺ is in fact close to that derived using the experimental 10Dq value. The difference, ΔR , between the R values for RbCaF₃:Mn²⁺ and RbCdF₃:Mn²⁺ becomes $\Delta R = (0.3\pm0.2)$ pm. This difference, although positive, appears to be a little smaller than $\Delta R = (1.8\pm1.2)$ pm derived from the experimental A_s values.

If we compare two systems such as for instance CsCaF₃:Mn²⁺ and RbCdF₃:Mn²⁺ it is clear from table 2 that $\Delta R < \Delta R_0$. To be more specific, in these cases it is found that $f \equiv \Delta R/\Delta R_0 \cong 0.35$. Thus, accepting the same f value in the comparison between RbCaF₃:Mn²⁺ and RbCdF₃:Mn²⁺, a difference $\Delta R \cong 1$ pm could be expected.

It is worth noting that as 10Dq for systems with MnF₆⁴⁻ units is measured mainly through the difference $\Delta = E_3 - E_1$, the dispersion of data for E_3 is also reflected in the 10Dq values. This situation could explain the slightly different R values obtained through equations (1) and (2). In the less favourable case of CsCaF₃:Mn²⁺, this difference is 0.9 pm.

Let us now focus on phase transitions of RbCaF₃ detected using optical parameters of the Mn²⁺ probe. Interesting results are obtained plotting the experimental first-order moment, M_1 , of the emission band associated with RbCaF₃:Mn²⁺ and comparing it to the corresponding variation displayed by CsCaF₃:Mn²⁺. Such results are given in figure 2. It can be seen that above 100 K the first moment M_1 undergoes in both cases a progressive increase upon heating. This increase as previously discussed [29] partially reflects the effects of thermal expansion giving rise to a decrement of 10Dq and thus an increase of the ${}^6A_{1g}(S) \rightarrow {}^4T_{1g}(G)$ energy. Nevertheless, at variance with what is observed in CsCaF₃:Mn²⁺ where the host lattice does not undergo a phase transition below 300 K [20], a slight but sensible change in the slope of $M_1(T)$ is detected in RbCaF₃:Mn²⁺. Therefore this anomaly can reasonably be related to the $O_h^1 \rightarrow D_{4h}^{18}$ structural phase transition experienced by the host lattice at 193 K. It is worth stressing however that the detection of such a phase transition through $M_1(T)$ requires a lot of measurements of M_1 in the 300-100 K range. In the present work M_1 has been measured for about 40 different temperatures in this range. As a result of the change of the slope at 193 K, the extrapolated value M_1 at 0 K (called $M_1(0)$) would be about 50 cm⁻¹ smaller for the tetragonal phase than for the cubic one. This can be related to a slightly higher 10Dq value for the former than for the latter case. If $\delta E_{CT}(0) = R_C(0) - R_T(0)$, where $R_{\rm C}(0)$ and $R_{\rm T}(0)$ denote the extrapolated Mn²⁺-F⁻ distances at 0 K for the cubic and the tetragonal phases, respectively, a value $\delta E_{CT}(0) \simeq 0.2$ pm is estimated using equation (2).

Upon lowering the temperature another anomaly is detected in the neighbourhood of 40 K (figure 2). Here M_1 experiences a small but abrupt blue shift of $\sim 100~\rm cm^{-1}$ which can in fact be related to the existence of another phase transition leading to a decrement of 10Dq upon cooling. Such a decrease would imply an *increase* of the average $\rm Mn^{2+}-F^{-}$ distance, R, equal to $\sim 0.5~\rm pm$ following equation (2), leading to a relative increase $\Delta R/R \simeq 0.2\%$. It is worth stressing now that the compound KMnF₃, apart from the $\rm O_h^1 \rightarrow \rm D_{4h}^{18}$ structural phase transition at $T_{c1}=184~\rm K$, experiences also the $\rm D_{4h}^{18} \rightarrow \rm D_{4h}^{5}$ phase transition at $T_{c2}=81.5~\rm K$. Dilatometric measurements [33] reveal that when this first-order transition takes place the length, L, of the sample undergoes an abrupt *increase*, $\Delta L/L$ being equal to 0.18%. This value is similar to that derived for the phase transition of RbCaF₃ at $T_{c2} \simeq 40~\rm K$.

Figure 2 depicts the temperature dependence of the luminescence lifetime for both $CsCaF_3:Mn^{2+}$ and $RbCaF_3:Mn^{2+}$ crystals. The experimental values for $CsCaF_3:Mn^{2+}$ have been fitted to the equation

$$\tau = \tau_0 \tanh(\hbar \omega_u / 2kT) \tag{3}$$

characteristic of phonon assisted electric dipole transitions. $\omega_{\rm u}/2\pi$ is the effective

phonon frequency. It can be seen that the fitting is very good with the exception of the experimental points lying in the T < 50 K region. The origin of this discrepancy, also seen in $\mathrm{Mn^{2+}}$ doped KMgF₃ and KZnF₃ [31], is at present not understood. In the case of RbCaF₃:Mn²⁺ a similar fitting of the experimental points in the 50-300 K range using equation (3) leads to a worse agreement. Nevertheless, this law is obeyed above 200 K in RbCaF₃:Mn²⁺ (solid line in figure 2). The progressive deviation of the experimental points below this temperature is also related to the existence of the $O_h^1 \to D_{h}^{18}$ structural phase transition even though this parameter is less sensitive than $M_1(T)$. Furthermore, the curve $\tau(T)$ does not provide any evidence on the existence of the phase transition at $T_{c2}=40$ K which is however well observed through M_1 . Though the origin of this different behaviour displayed by $M_1(T)$ and $\tau(T)$ with respect to the phase transitions of the host lattice is not well understood, it is worth noting that similar situations have been encountered for other materials containing $\mathrm{Mn^{2+}}$ [34, 35] or $\mathrm{Cr^{3+}}$ [36].

As a main conclusion, this work offers for the first time good evidence of two phase transitions in $RbCaF_3$ detected through an optical parameter of the Mn^{2+} probe. Also it provides a consistent view of the relation between optical and EPR parameters of Mn^{2+} in fluorides and the equilibrium Mn^{2+} -F⁻ distance, R. Further work devoted to clarifying the dependence of zero-phonon lines and Stokes shifts upon R for Mn^{2+} doped fluoroperovskites is now in progress.

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