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Understanding enormous redshifts in highly concentrated Mn²⁺ phosphors†

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Broad band near infrared (NIR) emission has recently been reported for a wide variety of concentrated Mn²⁺ phosphors. Typically, Mn²⁺ emits in the green to red spectral region, depending on local coordination. The enormous redshift to the NIR was explained by exchange coupling between Mn²⁺ neighbours at high Mn²⁺ dopant concentrations. However, the reported redshifts are an order of magnitude larger than expected for exchange coupling and also the absence of a shift in excitation spectra suggests that exchange coupling cannot explain the observations. Here, extensive concentration, temperature and time dependent luminescence studies are reported for $Mg_{1-x}Mn_xAl_2O_4$ (x = 0.01-0.5). The results show that the broad band NIR emission originates from NIR emitting trap centers, possibly Mn³⁺. High Mn²⁺ dopant concentrations enable efficient energy migration over the Mn²⁺ sublattice to these traps, consistent with the same excitation spectra for the green Mn²⁺ and NIR trap emission. Upon cooling to cryogenic temperatures energy migration is hampered and the green Mn²⁺ emission increases, especially in the most concentrated systems. Finally, the relative intensity of the NIR emission was varied by changing synthesis conditions providing further support that the NIR emission in concentrated Mn²⁺ phosphors originates from NIR emitting centers and not exchange coupled Mn²⁺ pairs.

Introduction

The development of new luminescent materials (phosphors) is crucial for improving white light LEDs for lighting and displays and also for enabling new applications. Important current topics in the LED phosphor industry concern narrow band red and green phosphors to improve the light quality, colour gamut and efficacy. 1-5 New applications are emerging and include human centric lighting and lighting for horticulture. 6-11 An interesting spectral region is the near infrared (NIR). In addition to applications in horticulture (controlling plant growth) and human centric lighting (adding NIR to the lamp spectrum for health benefits), NIR emitters are now gaining attention for chemical sensing. 12,13 For both narrow band phosphors and NIR emitters there is increasing interest in phosphors relying on the luminescence of the Mn²⁺ ion.^{14,15}

Mn2+ has a long history of applications in a variety of phosphors for lighting and displays. 16,17 The luminescence properties of Mn2+ are versatile and can be understood based on the Tanabe-Sugano diagram for the 3d⁵ configuration. Mn²⁺ typically yields narrow band green emission in tetrahedral coordination and red emission in octahedral coordination

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resulting from the ${}^{4}T_{1} \rightarrow {}^{6}A_{1}$ transition. The luminescence quenching temperature can be very high, making Mn2+ phosphors attractive for applications where high temperatures are reached, e.g. in white light LEDs (wLEDs). 18 Drawbacks are the weak absorption (ε < 1 M⁻¹ cm⁻¹) and long luminescence life time ($\tau > ms$) as all transitions within the 3d⁵ configuration are spin- and parity forbidden. The weak absorption can be mitigated by combining Mn2+ with a strongly absorbing sensitizer and the emission life time can be shortened by incorporation in a host lattice with heavier elements (stronger spin-orbit coupling) or through magnetic interactions, e.g. in Mn²⁺ pairs (partially lifting the spin selection rule). 19-25 In the past decade a new emission was reported for Mn2+: NIR emission from exchange coupled Mn²⁺-pairs.²⁵⁻³⁴ This broad band NIR emission was observed in a variety highly doped Mn2+ phosphors and deemed promising for applications in both lighting and sensing.

Before considering these recent findings of NIR emission in highly-doped Mn²⁺ phosphors, it is good to discuss early work on the optical properties of exchange-coupled Mn²⁺ pairs. In an insightful review in 1988 McCarthy and Güdel summarize experimental findings and theoretical explanations for the role of exchange coupling in redshifting the Mn2+ emission and shortening the emission life time. 20 The spectral shifts are determined by the exchange coupling parameter J in the ground state and J_{ex} in the excited state. Depending on distance, geometry and bridging ligands, values for J and J_{ex} vary and are typically tens of cm⁻¹. The redshift and shortening of

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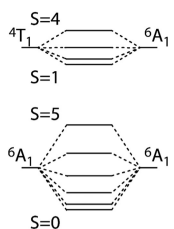


Fig. 1 Schematic of exchange coupling between Mn²⁺ neighbours in the 6 A₁ ground state (S = 5/2) giving rise to total spin S varying from 0 to 5 while exchange coupling with one ion in the ${}^{4}T_{1}$ excited state (S = 3/2) gives rise to total spin S varying from 1 to 4.20,21 In case of antiferromagnetic interaction S = 0 is the lowest state while for ferromagnetic interaction S = 5 will be the ground state for the exchange coupled pair. Note that the actual exchange splitting is much smaller than the splitting in the schematic figure.

decay time are explained using an energy level diagram (Fig. 1) in which the total spin for the pair in the ground state can vary from 0 to 5 (with steps of 1) depending on the relative orientation of the S = 5/2 spins of the exchange coupled Mn^{2+} ions and of the S = 5/2 and S = 3/2 spins for the pair with one ion in the excited state. The total spin magnetic moments for the pair can thus be 0, 1, 2, 3, 4 or 5 in the ground state and 1, 2, 3 or 4 in the excited state, giving rise to partially spin-allowed transitions between pair states with the same total spin. In the review several examples of exchange coupled pairs are discussed, showing redshifts of typically 10-100 cm⁻¹, for example for Mn²⁺ pairs in CsMgCl₃, KMgF₃ and for Cr³⁺ pairs in ruby (Al₂O₃:Cr³⁺).^{20,21,36,37} For several commercial phosphors the concentration dependent redshift was later also experimentally observed, theoretically explained and related to phosphor performance. In the famous green willemite phosphor Zn₂SiO₄:Mn²⁺ the emission maximum shifts from 520 to 529 nm (\sim 250 cm⁻¹) upon raising the concentration from 0.5 to 20% Mn^{2+,38,39} In the electroluminescent ZnS:Mn2+ phosphor similar Mn-concentration dependent shifts are observed as well NIR emission. 40,41 The NIR emission was explained by energy migration to NIR emitting centers of which the nature was not known. In an overview of luminescence properties of exchange coupled pairs of transition metal (TM) ions in 2001 also Mn2+-Mn2+ pairs were discussed and a spectral shift from 519 to 528 nm accompanied by life time shortening from 7.2 to 4.5 ms was reported for MgAl₂O₄ doped with 0.2 to 10% Mn^{2+,21} Even though not all aspects of exchange coupling can be quantitatively explained, the influence of exchange coupling on the Mn²⁺ (and other TM ions) luminescence properties are well documented.

In the past decade unexpected NIR emission for heavily doped Mn²⁺ phosphors was reported for a wide variety of materials, including KZnF3, MgGa2O4, MnS, CaO, MgAl2O4, CsMnF3,

(Gd,Y)₃(Ga,Al)₅O₁₂ and Li₂ZnSiO₄. ²⁶⁻³⁵ In 2014 the first papers reported a strongly redshifted NIR emission band in addition to the 'normal' Mn²⁺ emission in KZnF₃ and MgGa₂O₄. ^{26,27} In KZnF₃ the normal 585 nm emission intensity decreased upon raising the Mn²⁺ concentration above 10% and a strong NIR emission at 770 nm appeared. Similar observations in MgGa₂O₄ revealed green emission shifting from 507 to 518 nm for Mn²⁺ concentrations of 1 to 20%. This green emission intensity strongly decreased upon raising the Mn2+ concentration and was accompanied by a strong increase of a broad 770 nm NIR emission band, gaining intensity for Mn²⁺ concentrations above 10%. In both cases the visible Mn^{2+} emission was assigned to the ${}^4T_1 \rightarrow$ ⁶A₁ transition on isolated Mn²⁺ while the NIR emission band was assigned to exchange coupled Mn2+ pairs, in the case of KZnF3 Mn²⁺ in adjacent octahedral sites and in the case of MgGa₂O₄ to Mn²⁺ ions in edge-sharing neighbouring tetrahedral and octahedral sites. The exchange coupling theory, as outlined in ref. 19, 20, 41 was cited to explain the redshift. In the years following many more examples were reported and included in reviews on Mn²⁺ doped phosphors. For example, in MnS deep red emission at 710 nm was assigned to isolated Mn2+ and two NIR 900 and 1380 nm emission bands were explained by emission from nextnearest and nearest neighbour exchange coupled pairs. 28 DFT calculations were done to support similar assignments in CaO:Mn²⁺.²⁹ In a recent paper a difference in thermal quenching behavior of the two emission bands in heavily Mn-doped Li₂Zn-SiO₄ was used for temperature sensing.³⁴

The vast number of papers and host lattices showing broad band NIR emission at high Mn²⁺ doping concentrations seems convincing but the redshifts reported for the exchange-coupled Mn²⁺ pairs are very large, varying from 3400 cm⁻¹ in (Gd,Y)₃ (Ga,Al)₅O₁₂ and Li₂ZnSiO₄ to 6800 cm⁻¹ in MgAl₂O₄ and MnS. 28,32 These shifts are more than ten times larger than those reported in earlier work, 20,21,36,37,40,42,43 where clear evidence was presented for the magnitude of the exchange splitting parameters J and J_{ex} based on temperature dependent luminescence and decay measurements as well as experiments in magnetic fields. In fact, the small shifts of the visible emission bands of Mn²⁺ upon raising the concentrations (e.g. from 507 to 518 nm in MgGa₂O₄) are consistent with earlier results on redshifts expected for exchange coupled Mn²⁺ pairs. In addition, no large shifts are observed in the excitation spectra which is unexpected for high concentrations of exchange coupled Mn-pairs with a strongly distorted energy level scheme that would be expected based on the enormous redshifts reported for the emission. It is the aim of this work resolve the discrepancies between the recent papers and earlier work on exchange coupled Mn-pairs.

To find the origin for the NIR emission in heavily Mn²⁺ doped phosphors, here the luminescence properties of Mg_{1-x}- Al_2O_4 :Mn_x (x = 0.01 to 0.5) are investigated in detail. The MgAl₂O₄:Mn²⁺ phosphor is well studied and serves as a model system to provide insight. As Mn²⁺ doping is raised, a prominent NIR emission appears, consistent with the results reported in. 32 However at low temperatures (<100 K), this NIR emission decreases, while the typical green emission of $Mn^{2+}(^{4}T_{1} \rightarrow {}^{6}A_{1})$ reappears. In addition, the relative intensity of the NIR

emission is shown to depend on synthesis conditions and host stoichiometry. Based on our experimental findings, we propose a different origin for the NIR emission: it does not originate from exchange coupled Mn²⁺ pairs but from NIR emitting traps. At high doping concentrations efficient energy migration between excited Mn²⁺ ions facilitates energy transfer to this NIR luminescent center, which we tentatively assign to trace amounts of Mn³⁺. Energy migration is hampered at cryogenic temperatures, explaining the reappearance of the green Mn²⁺ emission. Synthesis conditions favoring incorporation of Mn3+ result in a higher NIR emission intensity and are consistent with our assignment.

Methods

For the synthesis of microcrystalline Mg_{1-x}Mn_xAl₂O₄ the starting materials MgO (99.99%, Sigma Aldrich), Al₂O₃ (99.99%, chempur), and MnCO₃ (99.99%, Sigma Aldrich) were mixed in stoichiometric amounts in an agate mortar and pestle. The mixture was wetted with a few drops of acetone (to prevent microparticle aggregation), and ground manually for 15 minutes. The mixtures were then annealed twice in a tube oven for a total of 18 h at 1500 °C in a reducing atmosphere (H₂/N₂ 20/80) and were reground in between.

The powders were examined using powder X-ray powder diffraction for phase purity. A Phillips PW1729 X-ray generator, Cu Ka source was used at 40 kV operating voltage and 20 mA current. The step size resolution was 0.02° 2θ . Photoluminescence (PL) spectroscopy was performed using an Edinburgh Instruments FLS-920 fluorescence spectrometer. The photoluminescence (PL) and PL excitation (PLE) measurements were recorded using a 450 W Xe lamp as excitation source and a Hamamatsu R928 PMT or a Hamamatsu N7422 PMT detector for the visible and NIR spectral region, respectively. To properly compare the intensity of the NIR with the VIS detector the spectra were normalised on the 650-700 nm peaks that were measured with both detectors. Spectra are corrected for the instrumental response. PL decay measurement were performed by using a tuneable optical paramagnetic oscillator (OPO) Opotek Opolette HE 355II pulsed laser (pulse width: 10 ns; repetition rate: 20 Hz) as excitation source and the Hamamatsu R928 PMT combined with the Multi Channel Scaling (MCS) time resolved measuring card in the Edinburgh spectrometer (10 µs time resolution). For temperature dependent measurements a liquid He cooled cryostat from Oxford Instruments was used.

Results and discussion

Concentration dependent luminescence

To investigate the concentration dependent luminescence properties of Mg_{1-x}Mn_xAl₂O₄, a series of samples was made under identical synthesis conditions with x = 0.01, 0.02, 0.05,0.1, 0.2 and 0.5. The materials came out of the tube oven as white powder for low-doped samples while at high doping concentrations (x > 0.10) the body color was slightly yellowish. XRD powder diffraction confirmed that all samples were phase pure (ESI† Fig. S1). The diffraction angles showed a continuous decrease for increasing Mn content as expected based on

Vegard's law and the fact that the ionic radius of Mn²⁺ is larger than that of Mg²⁺ and thus the crystal unit cell expands upon increasing Mn-content. Note that in the MgAl₂O₄ host it is expected that Mn²⁺ mainly occupies the Mg²⁺ site (similar size, same charge) creating a tetrahedrally coordinated Mn. 44 Mn2+ emission has also been reported for Mn²⁺ on the Al³⁺ site in $Mg_{1-x}Mn_xAl_2O_4$.⁴⁵

To characterize the Mn²⁺ luminescence for isolated Mn²⁺ ions, first PL and PLE spectra were recorded for MgAl₂O₄ doped with 1% Mn. In Fig. 2 the PL spectrum is shown. Narrow band green emission peaking at 520 nm is observed, consistent with earlier reports. The emission band is assigned to the ${}^4T_1 \rightarrow {}^6A_1$ transition on Mn²⁺. In addition there is a very weak sharp emission line at 680 nm. This is typical for Cr³⁺ at the Al³⁺ site and careful measurement of the position of the ${}^{2}E \rightarrow {}^{4}A_{2}$ zerophonon line and vibronic structure show that these are identical to what has been reported for Cr3+ in MgAl2O4 providing evidence for the incorporation of a small amount of Cr³⁺ in our materials. 46 Cr is a common contaminant in the Al₂O₃ precursor. The recorded PLE spectrum of the green Mn²⁺ emission shows six distinct peaks, at 280, 350, 380, 425, 450 and 480 nm. These are all d-d transitions of the Mn2+ and can be assigned to transitions from the ${}^{6}A_{1}$ ground state to the ${}^{4}A_{2}({}^{4}F)$, ${}^{4}E({}^{4}D)$, $^4\text{T}_2(^4\text{D})$, $^4\text{A}_1/\text{E}(^4\text{G})$, $^4\text{T}_2(^4\text{G})$ and $^4\text{T}_1(^4\text{G})$ excited states, respectively. No broad emission around 650 nm is observed. Emission at this wavelength is typical for Mn²⁺ at the octahedral Al site.⁴⁵

The concentration dependence of the luminescence was investigated by measuring luminescence spectra for all Mn²⁺ concentrations over a wide spectral range, extending to 1200 nm. Note that in the past NIR emissions from highly Mn-doped samples may have been missed because of low sensitivity of detectors (photomultipliers) in the NIR. In Fig. 3 the emission spectra are shown. The green 520 nm emission band increases in intensity up to 5% Mn²⁺ followed by a decrease above 10%. For the highest concentrations, 20 and 50% Mn²⁺, the green emission is almost completely quenched. The emission maximum shifts from 518 to 526 nm upon raising the Mn²⁺ concentration from 1 to 50%. In the concentration region where

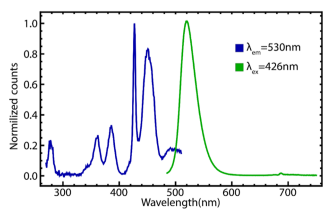


Fig. 2 Photoluminescence (green) and excitation (blue) spectrum of Mg_{0.99}Mn_{0.01}Al₂O₄ at 300 K. The PL spectrum was recorded under 426 nm excitation, the PLE spectrum for 530 nm emission.

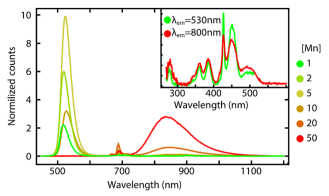


Fig. 3 Emission spectra of $Mg_{1-x}Mn_xAl_2O_4$ for x = 0.01,0.02,0.05,-0.10,0.20,0.50, excited at 425 nm. From 450 to 800 nm the spectra were measured with a R928 detector and in the NIR region (600 to 1200 nm) with a N7422 detector. The peak at \sim 680 nm was used to scale the spectra. The inset shows the excitation spectrum of the green emission of the 1% Mn and of the NIR emission of the 50% Mn material.

the green emission is quenched, a NIR emission band around 830 nm is observed to increase with Mn-dopant concentration. The highest NIR intensity is found for the 50% dopant concentration. The present results are in good agreement with the trends observed for the visible and NIR emission in other host lattices where the NIR emission was assigned to exchange coupled Mnpairs. Specifically, the spectra and relative intensities of visible and NIR emission are also consistent with results reported for concentration dependent Mn²⁺ luminescence in MgAl₂O₄. ^{21,32}

In the inset of Fig. 3, the excitation spectra of the green emission for the 1% Mn and the NIR emission in 50% doped MgAl₂O₄ are shown. The excitation spectra are similar and the assignment to Mn²⁺ d-d transition as discussed above can explain the features. The peak positions are almost identical while the excitation bands are broadened for the 50% doped host. The broadening can be explained by inhomogeneous broadening. The disorder in the Mg/Mn sublattice is maximized in the 50/ 50% material and will give rise to variations in the local crystal field splitting, induced by the difference in ionic radii for Mn and Mg. Variations in the local Mn-Mg distribution over nearest and possibly next-nearest neighbour lattice sites will thus give rise in differences in the crystal field splitting for Mn²⁺ and disorder is maximized in the host lattice with 50% Mn²⁺.

To further investigate the origin of the NIR emission band, luminescence decay curves were recorded for both the green and NIR emission as a function of concentration following pulsed excitation (~10 ns pulse duration) at 420 nm. Excited state dynamics can give valuable information on the role of energy transfer processes. The results in Fig. 4(a) for the green emission show that for 1 and 2% Mn the decay curves are close to single exponential with a decay time of 6.1 ms for 1 and 2% and drops to 0.5 ms at 20% Mn. The 6.1 ms is in agreement with decay times reported for 1-2% Mn2+-doped samples in ref. 21 For concentrations of 5% and higher the decay becomes faster and increasingly non-exponential which can be explained by shortening of the life time by exchange coupling and, especially for higher Mn2+ concentrations, energy migration and transfer to trap centers. The decay curves for the NIR emission in Fig. 4(b) also show decay on a ms time scale. The decay curves are non-exponential and decay times decrease for higher Mn²⁺ concentrations. The similar decay behavior observed for the NIR and green emission (non-exponential with an average decay time in the ms range) can be expected when the trap emission is populated by transfer from Mn²⁺ ions and the decay profile of the trap emission reflects the decrease in population of Mn²⁺ that feeds the trap states.

Based on the concentration dependent luminescence measurements an alternative explanation for the NIR emission band can be given. Upon increasing dopant concentrations above 10% resonant energy transfer between Mn²⁺ dopant ions leads to concentration quenching. Due to the efficient energy transfer between neighbouring Mn2+ ions the excited state probes a larger volume and even in case of low concentrations of trapping sites, efficient energy transfer to trap states can occur. Especially above the percolation point (the concentration at which a 3D connected network of dopants is formed) efficient energy migration to traps occurs, leading to quenching of the dopant emission and, in the case of luminescent traps, a strong increase in trap emission. This phenomenon is well-known in the field of luminescent materials. 47-49 The concentration dependent luminescence properties for MgAl2O4:Mn are consistent with this model: above 10% Mn²⁺ the NIR trap emission intensity rapidly increases while the green Mn²⁺ emission drops. The luminescence decay time of the Mn2+ donor decreases by migration mediated energy transfer to the trap states. The excitation spectrum for the

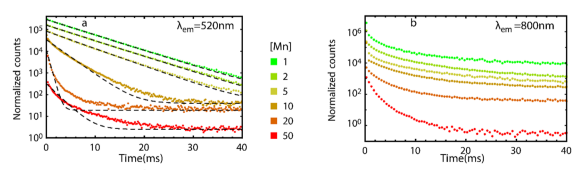


Fig. 4 Luminescence decay curves of Mn²⁺ in Mg_{1-x}Al₂O₄:Mn_x (x = 0.01, 0.02, 0.05, 0.10, 0.20, 0.50) for pulsed $\lambda_{ex} = 427$ nm excitation (curves are offset for clarity). (a) Decay curves for λ_{em} = 520 nm, the dashed lines show the fit assuming mono exponential decay. (b) Decay curves for λ_{em} = 830 nm.

trap emission is identical to that of the Mn2+ ions as the concentration of (and thus direct absorption by) the traps is very low. Following absorption by Mn²⁺ efficient transfer feeds the NIR emitting traps and this explains why the excitation spectrum for the NIR trap emission coincides with the Mn²⁺ absorption spectrum.

Note that in the earlier explanation (green emission from isolated Mn²⁺, NIR emission from exchange coupled Mn²⁺ pairs) there is no clear explanation for the fast drop in decay time for the green emission. Isolated Mn²⁺ is expected to have a long ms decay time while the exchange coupled Mn-pairs have a faster decay by partial lifting of the selection rules. Also the similarity in excitation spectra is unexpected for the model in which the NIR emission is explained by exchange coupled Mnpairs. Strong exchange coupling is required to explain the large redshift in emission and this large change in energy level structure should then also be reflected in changes in absorption transition energies. This is not observed. In other reports where strongly redshifted emission in concentrated Mn²⁺ phosphors is reported²⁶⁻³⁵ also no corresponding significant change in the excitation or absorption spectra is observed while it would be expected for such strong exchange coupling. Only for γ -MnS an additional 606 nm excitation band appeared in the excitation spectrum of the NIR but this can be due to direct excitation of the NIR emitting centers.²⁸

Temperature dependent luminescence

To further elucidate the nature of the NIR emission in highly doped Mn²⁺ luminescent materials, temperature dependent measurements can help. Energy migration involves many consecutive energy transfer steps between neighbouring Mn²⁺ ions. Small variations in local environment give rise to small energy differences between the ⁴T₁ excited state of different Mn²⁺ ions. At room temperature these small energy differences can be easily compensated by absorption or emission of phonons, making resonant energy transfer possible. However, upon cooling to cryogenic temperatures, the excitation energy will be trapped at Mn²⁺ ions which have their ⁴T₁ excited state at lower energies. Further transfer to a Mn2+ neighbour with a higher energy ⁴T₁ state would require absorption of a phonon which is not available as the phonons are frozen out a cryogenic temperatures. Extensive research on energy migration for especially lanthanide ions has beautifully demonstrated how at low temperatures energy migration is hampered and emission from perturbed lanthanide ions is observed as a slightly red shifted sharp line emission. 50,51 Interestingly, also for concentrated Mn²⁺ thermally activated energy migration to NIR emitting trap centers has been reported.⁵²

To investigate the role of energy migration in feeding the NIR emission in Mg_{1-x}Mn_xAl₂O₄ temperature dependent emission spectra down to cryogenic temperatures were performed for the MgAl₂O₄:50% Mn sample. The emission spectra are shown in Fig. 5. Upon cooling to 100 K there are no large changes but below 100 K a rapid increase in the green emission intensity observed while the NIR emission intensity decreases. The inset in Fig. 5 shows the integrated emission intensities of

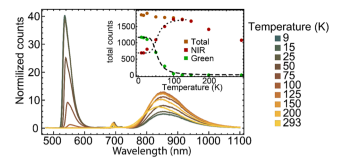


Fig. 5 Temperature dependent emission spectra of MgAl₂O₄:50% Mn, $\lambda_{\rm ex}$ = 426 nm. The inset shows the integrated intensity of the green and NIR emission bands. The dashed line shows the fit to determine the activation energy (ΔE of 200 cm⁻¹) for energy migration. The dotted line is a visual guide to show that the NIR emission increases simultaneously with the decrease of the green emission.

the NIR and visible emission bands as a function of temperature, normalized to the intensity at 4 K. The temperature dependence is consistent with efficient energy migration over the Mn²⁺ sublattice at room temperature, feeding the NIR emitting traps, which is hampered upon cooling below 100 K. The NIR emission dominates above 100 K and shows a small decrease upon raising the temperature to 300 K, possibly by thermal quenching of the NIR luminescence.

To quantify the activation energy that is necessary to overcome these energy fluctuations an Arrhenius equation was used to fit the observed temperature dependence of green emission intensity, as shown with the dashed line in Fig. 5. An effective ΔE of 200 cm⁻¹ was found. This value is consistent with what can be expected based on the broadening of the excitation bands for Mg_{0.5}Mn_{0.5}Al₂O₄ and also with earlier results by Güdel et al. on MnF2 single crystals where it was shown that perturbations by Ca impurities could lower the energy levels of nearby Mn²⁺ ions by 250 cm⁻¹.⁵³ In Fig. 6 the temperature dependent energy transfer is schematically depicted. At room temperature (Fig. 6(b)) efficient energy transfer between Mn2+ neighbours allows for rapid energy migration to NIR emitting trap centers resulting in dominant NIR emission for concentrations above the percolation point where an interconnected 3D network of Mn²⁺ ions is formed. At cryogenic temperatures (Fig. 6(a)) small energy differences between neighbouring Mn²⁺ ions hamper energy migration and result in a strong enhancement of the green Mn2+ emission.

The observed temperature dependence of the green and NIR emission is not consistent with the assignment of the green emission to isolated Mn2+ ions and the NIR emission to exchange coupled pairs. The concentration of isolated ions and exchange coupled pairs does not change with temperature and thus the strong increase of the green emission upon cooling from 100 K to 4 K cannot be explained by a model in which the green emission results from isolated ions and the NIR emission from exchange coupled pairs, especially for high Mn-doping concentrations where almost all Mn²⁺ ions are expected to be in pairs. A similar temperature dependence was previously even reported for fully concentrated Mn-system (e.g. MnS and CsMnF3) where

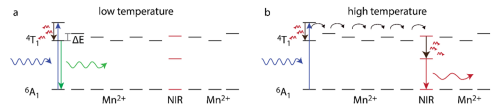


Fig. 6 Schematic illustration of energy migration to NIR emitting trap centers. (a) At low temperature energy migration is hampered by the lack of thermal energy to bridge energy differences leading to trapping the excitation energy at Mn²⁺ sites with slightly lower ⁴T₁ state resulting in an enhanced green Mn2+ emission and reduced NIR trap emission. (b) Energy migration to NIR emitting centers at room temperature where the small energy differences of the ⁴T₁ states for neighbouring Mn²⁺ ions can be thermally overcome by lattice vibrations leading to dominant NIR emission.

the isolated Mn²⁺ visible emission appeared upon cooling to 10 K. Clearly in a fully concentrated Mn2+ system all Mn2+ ions are in pairs and no isolated Mn²⁺ ions exist. For the CsMnF₃ material the observation of isolated Mn2+ emission at low temperature was attributed to competition between antiferromagnetic and ferromagnetic coupling interaction while for MnS no explanation was given for the observation of isolated Mn2+ visible emission upon cooling. In the MgAl₂O₄ system the NIR pair emission was assigned to pairs of Mn2+ ions on 'normal' four-coordinated Mg sites and six-coordinated Al3+ sites. Analysis of EXAFS data was used to determine that for 5% or less Mn²⁺ doping all Mn²⁺ was on IV-coordinated sites and for 10% or more $\mathrm{Mn}^{2+} \sim 8-10\%$ of Mn2+ ions occupied octahedral VI-coordinated sites. This can explain an increase of isolated Mn2+ emission upon cooling by hampering energy migration to Mn-pairs at cryogenic temperatures, similar to the energy migration to NIR trap centers discussed above. Unfortunately, the evidence for a significant Mn²⁺ concentration in octahedral sites is inconclusive. The EXAFS data show poor fits and most importantly, no red Mn²⁺ emission typical for Mn²⁺ on octahedral sites can be observed for any Mn²⁺ concentration.

The temperature dependence of luminescence decay curves for the green and NIR emission in MgAl₂O₄:50% Mn is shown in Fig. 7. At low temperatures (<50 K) the decay curves are characterized by a single exponential tail and a fast initial decay. This shape is characteristic of single step energy transfer and limited or no energy migration. One-step energy transfer to the NIR emitting centers gives rise to faster initial decay for those Mn²⁺ ions close to a NIR center and the single exponential tail (with decay times longer than 6 ms) reflects the radiative decay for Mn²⁺ ions which are not in proximity to a NIR emitting center. Above ~ 50 K the decay in the tail becomes faster indicating an onset of energy migration that becomes faster as the temperature is raised and

results in more efficient energy transfer to NIR centers through energy migration, depopulating the green emitting excited states. The temperature dependence of the NIR emission is more complex. It is on time scales similar to the decay of the green emission, indicating that the observed decay dynamics are controlled by the feeding states, in agreement with energy migration from Mn²⁺ to the NIR emitting centers.

Influence of non-stoichiometry

The results and discussion presented above provide strong evidence that the broad NIR emission originates from trap centers. Excitation of Mn²⁺ and energy migration over the Mn²⁺ sublattice is followed by efficient energy transfer to the NIR emitting center. A candidate for the NIR center is Mn3+. It is known that Mn3+ can show efficient NIR emission^{52,54-57} in the 700-1000 nm spectral region and the 3+ valence state is a well-known for Mn. The broad band Red/NIR emission from Mn^{3+} is typically assigned to the ${}^{1}T_{2} \rightarrow {}^{5}E$ emission but may also originate from the ${}^{5}T_{2} \rightarrow {}^{5}E$ transition. To investigate if the NIR emitter is Mn3+, non-stoichiometric samples doped with 10%Mn were synthesized while varying the amounts of MgO and Al₂O₃ precursor. It can be expected that excess Al₂O₃ will reduce the concentration of Mn³⁺ and that excess MgO will increase the Mn³⁺ concentration, based on the following equilibria reactions⁵⁸:

$$3MgO \iff 2Mg'_{Al} + Mg'_{Mg} + 3O'_{O} + V'_{O}^{\bullet \bullet}$$
$$3Al_{2}O_{3} + V'_{O}^{\bullet \bullet} \iff 4Al'_{Al} + 2Al'_{Mg} + 9O'_{O}$$
$$Mn'_{Mg} + V'_{O}^{\bullet \bullet} \iff Mn'_{Mg} + V'_{O}^{\bullet}$$

As Al is exclusively 3+ and Mg exclusively 2+, defect equilibria where is Mn is trivalent (either on a Mg²⁺ site or on a Al³⁺ site)

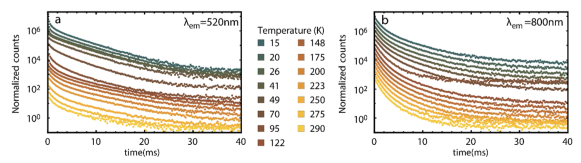


Fig. 7 Luminescence decay curves of the (a) 520 nm and (b) 800 nm emission in MgAl₂O₄:50% Mn following pulsed 427 nm excitation as a function of temperature.

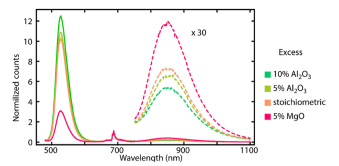


Fig. 8 Emission spectra of MgAl₂O₄:10% Mn synthesized with different excess of precursors. The NIR emission is also shown 30x enhanced for better visibility. The spectra were normalised at the 688 nm emission peak.

will be shifted towards lower Mn³⁺ concentration in the presence of excess Al3+ ions while excess Mg2+ will shift the equilibria towards more Mn³⁺. The emission spectra in Fig. 8 show indeed a strong increase in the NIR emission for 5% excess MgO and a decrease for 5 or 10% excess Al₂O₃.

The assignment of the NIR emission to trivalent Mn³⁺ is in agreement with the experimental results. At higher dopant concentrations the strong increase in NIR emission is not only caused by enhanced energy migration over the Mn2+ sublattice but also, for a constant fraction of Mn being in the 3+ state, the absolute concentration of NIR emitting centers would increase at higher dopant concentrations giving rise to a sharp increase in NIR emission intensity for higher Mn-concentrations. Still, the identification of the NIR emitting trap as Mn³⁺ should be taken with some caution. NIR emission is also observed in undoped MgAl₂O₄⁵⁹ this suggests that the NIR emission could also be caused by an intrinsic defect. In a recent paper by Sun et al. NIR emission from Mn²⁺ was observed and in line with the present findings not assigned to exchange coupled pairs but to Mn²⁺ in dodecahedral sites.60 It is notoriously difficult to unravel the nature of defects in luminescent materials and further research is needed to confirm the nature of the NIR emitting trap center for the many phosphors in which broad band NIR emission is observed at high Mn-doping concentrations and the nature of the NIR emitting trap center can vary for the different concentrated Mn2+ phosphors.

Discussion

The results and analysis presented above provide strong evidence that the enormous redshifts observed in concentrated Mn²⁺ phosphors do not originate from exchange coupled Mn²⁺ pairs. It is however important to realize that exchange interaction does affect the luminescence of transition metal ions, especially for spin-forbidden transitions. For Mn²⁺ much higher absorption strengths have been reported for Mn2+ pairs e.g. in KMgF3, $\text{CsMg}_{0.95}\text{Mn}_{0.05}\text{Cl}_3$ and $\text{CsMg}_{0.80}\text{Mn}_{0.20}\text{Br}_3.^{61,62}$ In excitation and absorption spectra additional strong exchange coupled Mn²⁺-pair bands appeared between 10 and 20 K as the S=1 pair state was thermally populated (see Fig. 1) and the partially spin-allowed transition to the S = 1 excited became possible. The temperature

range (10-20 K) and spectral shift of the pair absorption lines show that the exchange coupling strengths for Mn2+ are of the order of tens of cm⁻¹, consistent with theoretical calculations. In recent publications redshifted NIR emission was also reported for exchange coupled Cr3+ dimers with shifts of about 1000 cm-1 from the isolated $^{2}E \rightarrow {}^{4}A_{2} \text{ Cr}^{3+}$ emission around 700 nm. 63,64 These shifts are in line with redshifts expected for exchange coupled Cr³⁺ pairs. Exchange interaction in dimers is known to be stronger for Cr3+ than for Mn2+ which results in higher magnetic transition temperatures (e.g. $T_N \sim 308$ K for $Cr_2O_3^{65}$ vs. 118 K for MnO⁶⁶) and spectral shifts observed for exchange coupled pairs of Cr³⁺ ions are much larger than for Mn²⁺. In the review by McCarthy and Güdel examples, including Cr3+ pairs in ruby, are discussed with spectral shifts of hundreds of cm⁻¹ of the R-line emission for Cr³⁺ pairs.²⁰

Exchange interaction in transition metal ions pairs can give rise to stronger absorption and shorter emission decay times by relaxing the spin-selection rule and spectral shifts that vary depending on host lattice and transition metal ion. This can be exploited in the emerging field of new phosphors for NIR LEDs. The reported enormous redshifts for Mn²⁺ exchange coupled pairs of up to 7000 cm⁻¹ are however more than an order of magnitude larger than what can be expected for Mn2+ pairs and this, together with the absence of changes in the excitation spectra of the redshifted emission and the observation of a strong increase of normal Mn²⁺ emission in highly concentrated Mn²⁺ phosphors at cryogenic temperatures, warrants the conclusion that the enormous redshifts observed in the emission of concentrated Mn2+ phosphors does not originate from exchange coupled Mn2+ pairs but from NIR emitting trap centers.

Conclusions

The origin of the NIR emission in highly Mn-doped phosphors was investigated. Earlier reports assign the NIR emission to exchange coupled Mn2+-pairs in variety of host lattices. The reported spectral shifts are however more than 10 times larger than what can be expected based on exchange coupling parameters for Mn²⁺. Careful studies over a wide Mn-concentration range in $Mg_{1-x}Mn_xAl_2O_4$ (x = 0.01-0.5) indicate that energy migration towards NIR emitting trap centers is responsible for the NIR emission and not exchange coupled Mn-pairs. Temperature dependent measurement on MgAl₂O₄:50% Mn reveal a strong increase in the green emission intensity upon cooling to cryogenic temperatures, while the NIR emission intensity drops. This is consistent with energy migration that is hampered upon cooling as it requires thermal activation. The temperature dependence cannot be explained by a model in which exchange coupled Mn-pairs cause the NIR emission band in highly concentrated Mn-systems. Moreover, no redshifted absorption band is observed which would be expected for exchange coupled Mn-pairs. The NIR emitting center is tentatively assigned to Mn3+ and this assignment is supported by variations in the relative intensity of the NIR emission band in non-stoichiometric materials synthesized with excess MgO or Al₂O₃. Although the exact nature of the NIR emitting center is difficult to pinpoint it is evident that exchange coupled Mn-pairs cannot be responsible for the strongly redshifted emission in highly Mn-doped phosphors but that energy transfer to NIR emitting defects or impurities causes the NIR emission.

Conflicts of interest

There are no conflicts to declare.

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