Enhanced Red Phosphorescence in MgGeO₃:Mn²⁺ by Addition of Yb³⁺ Ions

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Enhancement of the T₂₁(G) → A₁₄(S) red emission of MgGeO₃:1% Mn²⁺ with addition of YbO₂ is reported. The initial phosphorescence in the YbO₂ doped sample is an order of magnitude stronger than that of the YbO₂-free sample. Thermoluminescence spectra indicate that the introduction of Yb³⁺ into the host produces traps with a suitable depth, resulting in the enhanced red phosphorescence of Mn²⁺. The possible mechanism for this phenomenon is explained by a competitive trapping model.

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Long-lasting phosphorescence (LLP) materials have long been of interest for various displays and signing applications. Blue and green oxide persistent phosphors, such as Sr₂Al₂O₆: Dy,Eu, SrAl₂O₄: Dy,Eu, exhibiting high brightness of phosphorescence with better chemical stability over sulfides are commercially available. However, commercial red-emitting oxide-persistent phosphors with a long persistence time have not yet been obtained.

Manganese ions (3d⁵ configuration) should be expected as an efficient activator for red long persistent phosphors due to their highly efficient luminescence and broad emission range from 490 to 750 nm in a different host. Many studies on the luminescence behaviors of the Mn²⁺ activator have been reported. The 3d™ multiplet energies of Mn²⁺ in crystals depend largely on the covalency interaction with the host crystal or the crystal field, because the 3d electrons of the transition metal ions are the outermost ones. The octahedral coordinated Mn²⁺ ion exhibits orange to red emissions.

Generally, either intrinsic or foreign defects are harmful to luminescent efficiency of activators because they could compete with activators and trap the excitation energy through nonradiative transitions. The density of traps plays a key role in the persistent emission of phosphor materials. To improve the phosphorescence intensity and persistent time, the introduction of defects induced by adding auxiliary activators is needed. In this article, the red phosphorescence enhancement of MgGeO₃:Mn²⁺ by doping Yb³⁺ into the phosphor is reported. The initial phosphorescence intensity in the doped sample is 1 order of magnitude higher than that of the Yb³⁺-free sample, and the persistent time is prolonged from several minutes to 30 min.

Experimental

Sample preparation.—The Mn²⁺-doped MgGeO₃ phosphors were prepared using the traditional solid-state reaction method. Analytical grade GeO₂, MgO, C₄H₆MnO₄·4H₂O, and Yb₂O₃ were used as starting materials. Those mixed-powder materials were homogenized, thoroughly ground with an agate pestle in an agate mortar, and then sintered at 1250°C for 3 h using alumina crucibles with alumina lids in air atmosphere.

Analysis and measurements.—The X-ray diffraction (XRD) data were collected by a Rigaku D/max-IB X-ray powder diffractometer using Cu Kα1 (λ = 0.15405 nm) radiation. The energy-dispersive X-ray spectrum (EDX) result was observed by a HITACHI-4800 scanning electron microscope equipped with an EDX.

The photoluminescence (PL) and excitation spectra as well as phosphorescence decay curves were measured on a Hitachi F-4500 fluorescence spectrophotometer equipped with a 150 W Xenon lamp as the excitation source. The phosphorescence was measured after irradiation by 254 nm UV light for 5 min. The thermoluminescence (TL) spectra were recorded by heating the irradiated samples from room temperature to 250°C using the EL-427A TL meter (Beijing Nuclear Instrument Factory) with a fixed heating rate at 2°C/s. All the measurements were carried out at room temperature except for the TL spectra.

Results and Discussion

Phase characterization.—The XRD patterns of MgGeO₃:1% Mn²⁺ singly doped and MgGeO₃:Mg²⁺:Yb³⁺ doped samples are shown in Fig. 1a. It is clearly indicated that the samples are chemically and structurally single-phased MgGeO₃, which corresponds to the JCPDS (no. 34-0281). There are no extra peaks observed in the XRD pattern from the YbO₂-added sample, suggesting that Yb³⁺ is well incorporated into MgGeO₃ lattice. The ionic radii of Yb³⁺ (r = 0.093 nm) is closer to that of Mg²⁺ (r = 0.072 nm) than that of Ge⁴⁺ (r = 0.053 nm); thus, we believe that Yb³⁺ ions prefer to occupy the Mg²⁺ sites. The inset of Fig. 1a shows a blueshift of the XRD peaks in the Yb doped sample compared to the Yb-free sample, which can be attributed to the lattice expansion because the ionic radii of Yb³⁺ is larger than that of Mg²⁺. Therefore, the blueshift provides the evidence of the Yb³⁺ incorporation into MgGeO₃ lattices, and the Yb³⁺ should substitute for the Mg²⁺. The EDX result as shown in Fig. 1b also confirms the presence of Yb in the MgGeO₃:Mn²⁺:Yb³⁺ doped sample, which indicate Yb³⁺ ions are incorporated into MgGeO₃:Mn²⁺:Yb³⁺ more clearly.

PL properties of MgGeO₃:Mn²⁺:Yb³⁺ phosphor.—The excitation and emission spectra of MgGeO₃:0.01 Mn²⁺ are presented in Fig. 2a. The emission spectrum exhibits a red emission band at 650 nm, originating from T₂₁(G) → A₁₄(S) transition of Mn²⁺ with an excitation at 254 nm. A blue emission band centered at 400 nm is attributed to the host. The excitation spectrum monitoring the 650 nm emission consists of a broadband centered at 254 nm and two shoulder bands centered at 209 and 238 nm, respectively. The band centered at 254 nm is due to the transition of Mn²⁺[A₁₄(S) → A₁₄(F)], and the two shoulders are related to the absorption of the host. In order to explain the emission and excitations related to the host, an undoped MgGeO₃ sample was also synthesized. As shown in Fig. 2b, two peaks centered at 209 and 238 nm,
respectively, are presented in the excitation spectrum and a broad-band emission peaked at 400 nm on the excitations is observed. The results indeed indicate that 209 and 238 nm excitations belong to the host absorption, and therefore, the MgGeO$_3$ is a self-activated material.

In addition, the excitation at 254 nm for the Mn$^{2+}$ doped sample overlaps with the host emission band. An intensity decrease of the host emission at 400 nm on 238 nm excitation is observed compared to that of the Mn$^{2+}$-doped sample, suggesting a possible energy transfer from the host to the Mn$^{2+}$ ions.

**Effect of Yb$_2$O$_3$ on the phosphorescence properties of MgGeO$_3$:Mn$^{2+}$.**—After removal of excitation source, the MgGeO$_3$:Mn$^{2+}$ sample yields weak red LLP. The introduction of Yb$_2$O$_3$ greatly improves the red LLP performance but weakens the PL intensity of Mn$^{2+}$. Figure 3 depicts the dependence of PL intensity (empty bars) and initial phosphorescence intensity (shaded bars) on Yb$^{3+}$ concentration in the codoped sample. The PL intensity in the samples decreases after Yb$_2$O$_3$ is incorporated into the host. The defects introduced by Yb$^{3+}$ quench the PL of Mn$^{2+}$. However, the defects act as traps to substantially improve the phosphorescence. As shown in Fig. 3, the phosphorescence from the Yb$_2$O$_3$ codoped samples appears more intense than that from the Yb$_2$O$_3$ free samples. The maximum of the phosphorescence intensity happens at the Yb$^{3+}$ concentration of 0.5%, which is 1 order of magnitude higher than that in the Yb$_2$O$_3$ free sample.

![Figure 1. Powder XRD patterns of MgGeO$_3$: (a) Mn$^{2+}$ and MgGeO$_3$:Mn$^{2+}$/Yb$^{3+}$ and (b) EDX for MgGeO$_3$:Mn$^{2+}$/Yb$^{3+}$.](image1)

![Figure 2. Excitation and emission spectra of (a) MgGeO$_3$:0.01 Mn$^{2+}$ and (b) undoped MgGeO$_3$ samples.](image2)

![Figure 3. Dependence of luminescence intensities (empty bars) and initial phosphorescence intensities (shaded bars) on Yb$^{3+}$ concentration in MgGeO$_3$:Mn$^{2+}$, xYb$^{3+}$ (x = 0, 0.1, 0.5, 1.0, 2.0, 5.0).](image3)
Figure 4 illustrates the time decay curve of phosphorescence in MgGeO$_2$:1% Mn$^{2+}$, 0.5 mol% Yb$^{3+}$ after irradiation with 254 nm UV light for 2 min. The persistent time is prolonged from several minutes to 30 min. The electron storage capacity of persistent phosphors is related to the trap depth and concentration. The Yb$^{3+}$-doped sample produces more defects that greatly enhance the phosphorescence. As shown in the inset of Fig. 4, the afterglow spectra are recorded at different times (2, 3, 4, and 5 min respectively) after the excitation source is switched off. It is found that the center (650 nm) and the profile of the red LLP emission remain unchanged and are consistent with PL emission under steady excitation. It is indicated that both LLP and PL result from the same $^{4}T_{1g} \rightarrow ^{6}A_{1g}$ transition of Mn$^{2+}$.

Figure 6. Possible process of the red LLP in the MgGeO$_2$:Mn$^{2+}$ phosphor. Possible mechanism of the LLP of MgGeO$_2$:Mn$^{2+}$, Yb$^{3+}$.—The LLP is due to the thermally stimulated recombination of holes and electrons that are trapped at the metastable states at room temperature. On the basis of the above TL curves and the PL spectra result, it should be thought that one energy transfer from the host to the Mn$^{2+}$ ions favors to create the LLP. It can be assumed that free electrons and holes in the sample are generated under UV excitation. The holes are captured by Mn$^{2+}$ ions, which form the excited state of Mn$^{4+}$ ions. The Mn$^{4+}$ substitution by Mn$^{2+}$ creates appropriate traps or oxygen vacancies associated with Ge$^{4+}$ ions, where electrons are captured. One part of the trapped electrons was released by heat at room temperature and turned back to the valence band, giving blue emission. This process has been approved by the PL of the undoped phosphor as shown in Fig. 2b. The majority of electrons stored in the electron traps transfers to the excited state of Mn$^{4+}$ ions. Finally, the excited state of Mn$^{4+}$ ions decay radiatively into the ground state with the red phosphorescence due to the $^{4}T_{1} \rightarrow ^{6}A_{1}$ transition resulting in red LLP of Mn$^{2+}$. The mechanism of the phosphorescence can be formulated in detail using a simplified scheme as shown in Fig. 6.

However, the electron traps depth is deepish because its relative temperature of TL peak is high at 133°C as shown in Fig. 5. The electrons are captured by the traps with deep depth, leading to poor performance of red LLP of Mn$^{2+}$ for several minutes. As the conclusion in the XRD results, when the trivalent Yb ions are codoped in the host matrix, the Yb$^{3+}$ ions tend to substitute for Mg$^{2+}$ ions rather than Ge$^{4+}$ because the ionic radii of Yb$^{3+}$ ($r = 0.093$ nm) is closer to that of Mg$^{2+}$ ($r = 0.072$ nm) than that of Ge$^{4+}$ ($r = 0.053$ nm). Yb$^{3+}$ ions replacing Mg$^{2+}$ ions produces Yb$_{Mg}$ defects with negative charge to maintain the electroneutrality of the phosphors. This negative Yb$_{Mg}$ site impurity traps with relative TL peak at 44°C, produced by codoped Yb$^{3+}$, is preferred as the electron trap with suitable depth. In the codoped sample, the amount of the defects located at 133°C decreases greatly, and most electrons are captured by the defects with suitable depth instead of the deeper defects, resulting in improvement of red LLP performance of Mn$^{2+}$.
Conclusion

Enhancement of red emission in MgGeO$_3$:Mn$^{3+}$ phosphor has been obtained with addition of Yb$_2$O$_3$. Influences of the excess Yb$^{3+}$ ions and introduced defects on red PL and LiP of the phosphor have been systematically investigated. The TL spectra prove that the introduction of Yb$^{3+}$ into the host produces traps with suitable depth, resulting in enhanced red phosphorescence of Mn$^{3+}$. The possible mechanisms of phosphorescence have been discussed.

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