Introduction

With the development of the economies of all countries in the world, environmental and energy issues have received increasing attention. The use of commercial greenhouses has grown rapidly in recent years to obtain green grains and high-quality vegetables. Light plays an important role in plant growth, because it can affect the growth rhythm of natural plants, such as delaying the plant blooming time or promoting plant blooming ahead of schedule. Phytochrome PR and PFR can influence the plant blooming time by converting to each other; PR needs more red light peaking at 660 nm to switch to the PFR biologically active state, and PFR captures more far-red light components. Therefore, research on far-red emitting phosphors for plant growth LEDs has also been made using the LLS:0.3%Mn4+,1.6%Mg2+ phosphors and a 365 nm emitting InGaN chip, which can emit far-red light that is visible to the naked eye. Importantly, the emission spectrum of the LLS:0.3%Mn4+,1.6%Mg2+ phosphors can match well with the absorption spectrum of phytochrome PFR, indicating the potential of these phosphors to be used in plant growth light-emitting diodes.

In this paper, we reported on the high-efficiency thermally-stable La2LiSbO6:Mn4+,Mg2+ (LLS:Mn4+,Mg2+) far-red emitting phosphors. Under 338 nm excitation, the composition-optimized LLS:0.3%Mn4+,1.6%Mg2+ phosphors which were made up of [SbO6], [LiO6], and [LaO8] polyhedrons, showed intense far-red emissions peaking at 712 nm (2Eg → 4A2g transition) with internal quantum efficiency as high as 92%. The LLS:0.3%Mn4+,1.6%Mg2+ phosphors also exhibited high thermal stability, and the emission intensity at 423 K only reduced by 42% compared with its initial value at 303 K. The far-red light-emitting device has also been made using the LLS:0.3%Mn4+,1.6%Mg2+ phosphors and a 365 nm emitting InGaN chip, which can emit far-red light that is visible to the naked eye. Importantly, the composition-optimized LLS:Mn4+,Mg2+ phosphors have excellent thermal stability with internal quantum efficiency (IQE) of the phosphors would have potential application in plant growth LEDs.

The phosphor-converted light-emitting diodes (LEDs), which show the advantages of energy-saving, long working lifetime, and environmental protection, have gained increasing attention as the next-generation light source. Research efforts have been devoted into developing high-efficiency red phosphors for white LEDs, such as NaMgGdTeO6:Mn4+, Na4MgZr(PO4)3:Eu3+, CaAl2O19:Mn4+, and Ca3MgV3O12:Eu3+. In sharp contrast, there is little research on the far-red emitting phosphors for plant growth LEDs. Thus, the research on the high-efficiency red phosphors for far-red LEDs used in plant cultivation has a far-reaching significance.

Mn4+ ions can be doped into the octahedral sites in the host as an activator, resulting in the efficient red or far-red emissions which are significant for plant growth. La2LiSbO6 compound with double-perovskite structure, which contains two octahedral structures, [SbO6] and [LiO6], have been reported by M. L. López et al. In this paper, La2LiSbO6:Mn4+,Mg2+ (LLS:Mn4+,Mg2+) far-red-emitting phosphors have been systematically studied. Under the excitation of 338 nm, LLS:0.3%Mn4+,1.6%Mg2+ phosphors exhibited intense far-red emission around 712 nm (2Eg → 4A2g transition), which matched well with the absorption spectrum of phytochrome PFR. Importantly, the internal quantum efficiency (IQE) of the LLS:0.3%Mn4+,1.6%Mg2+ phosphors reached as high as 92%. The temperature-dependent spectra illustrated that LLS:0.3%Mn4+,1.6%Mg2+ phosphors had excellent thermal stability with an activation energy of 0.12 eV, and the integral intensity at 423 K is 0.58-fold compared with its initial value at 303 K. All characteristics show that the LLS:xMn4+,yMg2+ far-red emitting phosphors would have potential application in plant growth LEDs.
Experimental section

LLS:xMn⁴⁺ (x = 0.1–1.5%) phosphors were prepared by solid-state reaction method. The amount of final product was fixed at 0.005 mol for each sample. The starting materials of Li₂CO₃ (analytical reagent, AR), Sb₂O₅ (99%), MnCO₃ (AR) and La₂O₃ (99.99%) were weighed according to the molar ratio of 1.03 : (1 – x) : 2x : 2 and ground in an agate mortar, then transferred to Al₂O₃ crucibles to pre-heat at 70 °C for 10 h in air. Then the obtained mixtures were ground thoroughly again, and calcined at 800 °C for 10 h in air. Subsequently, the obtained products were uniformly ground again and calcined at 1100 °C for 10 h in air to get the final samples. Finally, when the furnace was cooled down to room temperature, the obtained products were ground into fine powders for further characterizations. The LLS:0.3% Mn⁴⁺,yMg²⁺ phosphors were prepared by using the similar method, and the MgO (AR) was weighed at 0.8%, 1.2%, 1.6%, and 2.0% of the total molar amount of Li atom.

The X-ray diffraction (XRD) patterns of the phosphors were measured by using a Bruker D8 ADVANCE diffractometer with Cu Kα radiation (λ = 1.5406 Å). The morphology of the LLS:0.3% Mn⁴⁺ phosphors was recorded by using a field-emission scanning electron microscope (FE-SEM; TESCAN MAIA3). The photoluminescence (PL) spectra/PL excitation (PLE) spectra were measured by an Edinburgh FS5 spectrometer with a 150 W continuous-wave xenon lamp as the excitation source. The temperature-dependent spectra were recorded on the same spectrometer equipping with a temperature controller. The luminescence decay curves and the IQE of the as-prepared sample were measured by Edinburgh FSS spectrometer equipped with a pulsed xenon lamp as the excitation source and an integrating sphere coated with barium sulfate, respectively. The far-red light-emitting device was fabricated by coating the LLS:0.3%Mn⁴⁺,1.6%Mg²⁺ phosphors onto a 365 nm emitting InGaN chip, and the corresponding photoelectric properties of the device under a bias current of 300 mA were measured by using a spectroradiometer system (HAAS-2000, Everfine).

Results and discussion

Fig. 1(a) shows the XRD patterns of LLS:xMn⁴⁺,yMg²⁺ (x = 0.1%, 0.3%, 0.9% and 1.5%; y = 0.8% and 1.6%) phosphors. All the diffraction peaks coincided well with the stand card of La₂LiNbO₆ compound (JCPDS: 40-0895). Fig. 1(b) and (c) show the Rietveld refined results of LLS:0.3%Mn⁴⁺ phosphors and the crystal structure of the LLS:0.3%Mn⁴⁺ phosphors, respectively. From the Table 1 we could know that the LLS:0.3%Mn⁴⁺ phosphors belong to double perovskite structure with monoclinic crystal system and P121/n1 space group. The LLS:0.3%Mn⁴⁺ phosphors was build up with [Sb06], [Li06], and [LaO8], in which [Sb06] and [Li06] (see Fig. 1(c)) were bound together by sharing the same oxygen atoms (sharing corners with each other). The cell parameters of the LLS:0.3%Mn⁴⁺ phosphors were: a = 5.61710(8) Å, b = 5.73188(8) Å, c = 7.96305(11) Å, and V = 256.380(8) Å³. Considering the ion radii and coordination number (CN) of the Mn⁴⁺ ions (r = 0.53 Å, CN = 6), Sb⁵⁺ ions (r

![Fig. 1](image_url)
where 0.76 \( \equiv \frac{r}{b} \) and the ratio can be estimated by the following expression:

\[
D_t = \frac{R_m(CN) - R_d(CN)}{R_m(CN)} \times 100\%
\]

where \( R_m(CN) \) and \( R_d(CN) \) are the ionic radii of the host cations and the doped ions, respectively. Herein, \( R_d(CN) = 0.53 \) Å and CN = 6 for Mn\(^{4+}\) ions, \( R_m(CN) = 0.6 \) Å for Sb\(^{5+}\) ions, and \( R_m(CN) = 0.76 \) Å for Li\(^+\) ions. Thus, the \( D_t \) of Mn\(^{4+}\) ions substituted for Sb\(^{5+}\) and Li\(^+\) ions were 11.7% and 30.3%, respectively. Because the ratio of the Mn\(^{4+}\) ions substituted for Sb\(^{5+}\) ions was less than 30%, indicating in the LLS host the Mn\(^{4+}\) ions were substitute for Sb\(^{5+}\) sites in LLS host. Another parameter to determine which cation in the unit cell to be substituted by Mn\(^{4+}\) ions is the radii percentage difference \( D_r \) between the doping ions (here was Mn\(^{4+}\) ions) and the possible substituted ions (here were Sb\(^{5+}\) and Li\(^+\) ions), and the ratio can be estimated by the following expression:\(^{42,43}\)

\[
D_r = \frac{R(CN) - R_{\text{doped ions}}}{R(CN)} \times 100\%
\]

Fig. 3(b) presents the PL spectra of the LLS:0.3%Mn\(^{4+}\) (x = 0.1–1.5%) phosphors. The maximum emission intensity reached when x = 0.3%, indicating the existence of concentration quenching in LLS:0.3%Mn\(^{4+}\) phosphors. The critical distance \( (R_c) \) is a parameter for dopant ions in a particular lattice to determine whether exchange interaction or electric multipole interaction contribute to the concentration quenching, which can be obtained by the following formula:\(^{46,49}\)

\[
R_c = 2\left(\frac{3V}{4\pi N c^2}\right)^{1/3}
\]

where \( V \) is the volume of the unit cell and \( N \) is the number of cation in the unit cell to be substituted by Mn\(^{4+}\) ions; \( x_c \) refers to the optimal doping content of Mn\(^{4+}\) ions. Herein, \( V = 256.380(8) \) Å\(^3\), \( N = 2 \), and \( x_c = 0.3 \%). Thus, the value of \( x_c \) was calculated to be 43.4 Å (\( >5 \) Å), indicating that it was electric multipole interaction contribute to the concentration quenching in LLS:0.3%Mn\(^{4+}\) phosphors.\(^{49}\) Besides, the exact concentration quenching mechanism can be explored using the following formula expression:\(^{43,52}\)

\[
\frac{I}{I_0} = K(1 + \beta(x))^{n(x) - 1}
\]

where \( I \) and \( x \) represent the PL intensity and doping concentration of Mn\(^{4+}\) ions in LLS host, respectively; \( \beta \) and \( k \) are constants for specific host lattice; and \( \theta = 6, 8, \) and 10 stands for electric dipole–dipole, dipole–quadrupole and quadrupole–quadrupole interaction, respectively.\(^{53}\) Fig. 3(c) shows the relationship between log(\( x \)) and log(\( I/I_0 \)). We could know that the slope of the fitting line was \( -1.52 \), thus \( \theta = 4.56 \) which was close to 6, revealing that the quenching mechanism in LLS:0.3%Mn\(^{4+}\) phosphors is based on dipole–dipole interaction.

Fig. 3(d) depicts the luminescence decay curves of the Mn\(^{4+}\) ions in the LLS:xMn\(^{4+}\) (x = 0.1–1.5%) phosphors (\( \lambda_{\text{ex}} = 338 \) nm; \( \lambda_{\text{em}} = 712 \) nm). The decay curves of as-prepared samples could be well fitted by the following double exponential expression:\(^{44,56}\)

\[
I = A_1 \exp(-t/t_1) + A_2 \exp(-t/t_2)
\]

herein, \( I \) refer to the luminescent emission intensity at time \( t \), and \( A_1 \) and \( A_2 \) are constants; \( t_1 \) and \( t_2 \) are the lifetimes for the exponential component; the decay lifetimes were found to be 2.079, 2.047, 1.997, 1.943, 1.821, and 1.741 ms for \( x = 0.1 \%), 0.3 \%, 0.6 \%, 0.9 \%, 1.2 \%, and 1.5 \%, respectively. The decay lifetimes gradually decreased with the increasing Mn\(^{4+}\) concentration, which can be attributed to the enhanced non-radiative energy transfer of Mn\(^{4+}\) ions in LLS:xMn\(^{4+}\) phosphors.

To enhance the emission intensity and IQE of LLS:0.3%Mn\(^{4+}\) phosphors, a series of Mg\(^{2+}\) doped LLS:0.3%Mn\(^{4+}\) phosphors had been synthesized. Fig. 4(a) and (b) shows the PL spectra of LLS:0.3%Mn\(^{4+}\), yMg\(^{2+}\) phosphors doped with different Mn\(^{4+}\) concentrations and the corresponding relative intensity as a function of Mg\(^{2+}\) concentration, respectively. Clearly, the
Fig. 3  (a) The PLE and PL spectra of LLS:0.3%Mn4+ phosphors. (b) The PL spectra of LLS:xMn4+ (x = 0.1–1.5%) phosphors. (c) Plot of log(I/I₀) vs. log(x) of Mn4+ ions in LLS:xMn4+ phosphors excited at 338 nm. (d) The luminescence decay curves of LLS:xMn4+ (x = 0.1–1.5%) phosphors (λex = 338 nm; λem = 712 nm).

Fig. 4  (a) PL spectra of the LLS:0.3%Mn4+,yMg2+ phosphors excited at 338 nm. (b) The relative intensity as a function of Mg2+ concentration. (c) The excitation profile of the reference BaSO₄ and the emission spectrum of the LLS:0.3%Mn4+,1.6%Mg2+ phosphors at 338 nm excitation. (d) Decay curves of the LLS:0.3%Mn4+,yMg2+ (y = 0% and 1.6%) phosphors. (e) CIE chromaticity diagram of the LLS:0.3%Mn4+,1.6%Mg2+ phosphors. (f) PL spectrum of LLS:0.3%Mn4+,1.6%Mg2+ phosphors together with the absorption spectra of phytochrome Pr and Pfr.
emission intensity of LLS:0.3%Mn4+,1.6%Mg2+ phosphors was enhanced by the introduction of Mg2+ ions. Importantly, the emission intensity of the LLS:0.3%Mn4+,1.6%Mg2+ phosphors was about 2.5-fold of the LLS:0.3%Mn4+ counterpart. In the LLS:0.3%Mn4+,1.6%Mg2+ phosphors, the doped Mg2+ ions was served as the charge compensator. Because the radii of the Mg2+ ions (72 pm, CN = 6) and Li+ ions (76 pm, CN = 6) are relatively close, so the Mg2+ ions might substitute for Li+ sites or occupy Li+ vacancies, and then compensate the imbalanced charges in LLS:0.3%Mn4+ phosphors. As a result of charge compensation, the formation of the defects which served as quenching centers in the LLS:0.3%Mn4+ phosphors was avoided. Thus, the emission intensity and IQE of the LLS:0.3%Mn4+ phosphors were largely improved. Meanwhile, the IQE of the LLS:0.3%Mn4+ had also been largely improved by Mg2+ co-doping. The IQEs of LLS:0.3%Mn4+ and LLS:0.3%Mn4+,1.6%Mg2+ (see Fig. 4(c)) were measured to be 61% and 92%, respectively, which was higher than that of K2NaAlF6:Mn4+ (IQE = 58.4%), Ba3TiGeO6:Mn4+ (IQE = 35.6%), and Mg7Ga2GeO12:Mn4+ (IQE = 28.13%). The decay lifetime of the LLS:0.3%Mn4+,1.6%Mg2+ phosphors was longer than that of LLS:0.3%Mn4+ counterpart (see Fig. 4(d)). In addition, the CIE coordinates of LLS:0.3%Mn4+,1.6%Mg2+ phosphors were (0.733, 0.267), located in the far-red region. Importantly, as shown in Fig. 4(f), the PL spectrum of the LLS:0.3%Mn4+,1.6%Mg2+ phosphors matched well with the absorption spectrum of PFR, which indicated the LLS:0.3%Mn4+,1.6%Mg2+ phosphors had great potential to be used as far-red emitting materials in LEDs for plant growth application.

Fig. 5(a) shows the temperature-dependent spectra of the LLS:0.3%Mn4+,1.6%Mg2+ phosphors. The profiles of the emission spectra were almost the same, but the emission intensity gradually reduced with the increasing temperature. As shown in Fig. 5(b), the emission intensity of the LLS:0.3%Mn4+,1.6%Mg2+ phosphors at 423 K remained about 58% compared that at 303 K, which was better than that of Ca3La2W2O12:Mn4+ (29%), CaLaMgNbO6:Mn4+ (45%), SrLaScO4:Mn4+ (15%), and Gd2-ZnTiO3:Mn4+ (27.2%), indicating that the LLS:0.3%Mn4+,1.6%Mg2+ phosphors had good thermal stability. The corresponding activation energy can be obtained by using the following expression:

$$I(T) = I_0 \left[ 1 + c \exp \left( \frac{-\Delta E}{kT} \right) \right]^{-1}$$

where I(T) and I0 are the intensity at temperature T and the initial intensity, respectively; k is the Boltzmann constant and c is a constant. The activation energy of the LLS:0.3%Mn4+,1.6%Mg2+ phosphors, which was obtained from the fitting result in Fig. 5(c), was 0.12 eV. The possible thermal quenching behavior of the LLS:0.3%Mn4+,1.6%Mg2+ phosphors can be explained by a simple configuration coordinate diagram of 4A2g, 4T1g, 4T2g, and 2Eg levels, as shown in Fig. 5(d). At room temperature, the electrons on the ground state 4A2g were excited to the excited state under the excitation of 338 nm or 481 nm, and then return to the lowest excited state 2Eg by nonradiative transition. Eventually, the electrons on the 2Eg level return to 4A2g level with the emission of far-red light (the red line). But when the...
temperature increased, the electrons on the $^2E_g$ level are more likely to get enough energy to reach the crossover point of the states of $^4A_{2g}$ and $^2E_g$, then return to the ground state by non-radiative relaxation with no light emission, which result in the decrease of the emission intensity of the LLS:0.3% Mn$^{4+},1.6%Mg^{2+}$ phosphors.

In this host, the Mn$^{4+}$ ions were in a strong field environment, which can be reflected by the ratio of crystal filed strength ($D_q$) to Racah parameter $B$. The corresponding $D_q$ and $B$ can be determined by the following eqn (6) and (7):

$$D_q = \frac{E(4A_{2g} - 4T_{2g})}{10}$$

$$\frac{D_q}{B} = \frac{15(x - 8)}{(x^2 - 10x)}$$

where $x$ is defined as:

$$x = \frac{E(4A_{2g} - 4T_{1g}) - E(4A_{2g} - 4T_{2g})}{D_q}$$

And according to the fitting result of PL spectrum of the LLS:0.3%Mn$^{4+}$ phosphors, the peaking energy of $^2E_g \rightarrow ^4A_{2g}$ transition was about 14 045 cm$^{-1}$. Therefore, the Racah parameter $C$ can be calculated as following:

$$E(4T_{1g} \rightarrow 4A_{2g}) = \frac{3.05C}{B} + 7.9 - \frac{1.8B}{D_q}$$

Thus, based on the eqn (6)–(9), the values of crystal parameter $D_q$, $B$, and $C$ was 2079, 898, 2509 cm$^{-1}$, respectively. Meanwhile, the value of the $D_q/B$ is about 2.32, indicating the Mn$^{4+}$ ions were in a strong field environment in the LLS host. In a strong octahedral filed environment (surrounded by six oxygen atoms), as shown in Fig. 6(a), the original degenerate 3d orbitals of free Mn$^{4+}$ ions, which are $d_{x^2-y^2}, d_{z^2}, d_{xy}, d_{xz},$ and $d_{yz}$, split into two degenerate and three degenerate. The energy of the two degenerate consisting of $d_{x^2-y^2}$ and $d_{z^2}$ is higher than that of the three degenerate consisting of $d_{xy}, d_{xz},$ and $d_{yz}$. Fig. 6(b) also gives the simple schematic diagram of Mn$^{4+}$ ions energy level in strong octahedral filed environment. The possible transitions between different levels can be expressed as the following process. Under the excitation of 338 nm or 481 nm, the electrons on the ground state $^4A_{2g}$ were excited to the excited levels, and then relaxed the lowest excited level $^2E_g$. Finally, the far-red light peaking at 712 nm was emitted with the electrons on the $^2E_g$ level return to the $^4A_{2g}$ level.

A prototype far-red emitting LED device was fabricated by coating the as-prepared LLS:0.3%Mn$^{4+},1.6%Mg^{2+}$ phosphors on a 365 nm near-UV LED chip, and Fig. 7 displays the corresponding electroluminescence (EL) spectrum of the prototype LED device under a bias current of 300 mA. An intense far-red emission band in the wavelength rang of 650–780 nm ($^2E_g \rightarrow ^4A_{2g}$ transition) was observed and the far-red light around the device could be seen clearly by comparing the inset (i) and (ii) in Fig. 7. All the results indicate the LLS:0.3%Mn$^{4+},1.6%Mg^{2+}$ phosphors have a promising prospect in the plant lighting field.

**Conclusions**

In conclusion, double perovskite LLS:Mn$^{4+},Mg^{2+}$ far-red emitting phosphors with high IQE and excellent thermal stability were synthesized by traditional solid-state reaction method. The optimal doping concentration of Mn$^{4+}$ and Mg$^{2+}$ ions in
LLS: Mn$^{4+}$, Mg$^{2+}$ phosphors were 0.3% and 1.6%, respectively. Under the excitation of 338 nm, the LLS:0.3%Mn$^{4+}$,1.6%Mg$^{2+}$ phosphors showed an intense far-red-emitting band peaking at 712 nm and the emission intensity at 423 K remained 58% of its initial value at 303 K. By doping Mg$^{2+}$ ions in LLS:0.3%Mn$^{4+}$ phosphors, the luminescence intensity of Mn$^{4+}$ ions was enhanced by 2.5-fold and the corresponding IQE was also improved from 61% to 92%. Remarkably, the emission spectrum of the LLS:0.3%Mn$^{4+}$,1.6%Mg$^{2+}$ phosphors could match well with the absorption spectrum of the phytochrome Pfr. All the outstanding properties of the double perovskite LLS:Mn$^{4+}$,Mg$^{2+}$ phosphors make it be a promising far-red-emitting phosphor for applications in plant growth LEDs.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by the National Natural Science Foundation of China (No. 51502190), the Program for the Outstanding Innovative Teams of Higher Learning Institutions of Shanxi, and the Open Fund of the State Key Laboratory of Luminescent Materials and Devices (South China University of Technology, No. 2017-skllmd-01)

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