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Improving the luminescence properties of the near-infrared phosphor Ca_{0.8}Sr_{0.2}O:Eu²⁺ via energy transfer and its application in the concentration detection of ethanol solutions†

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In this work, a near-infrared (NIR) phosphor $Ca_{0.8}Sr_{0.2}O:Eu^{2+}$ was prepared via a solid-state method, and its luminescence properties were improved by co-doping with Ce^{3+} ions. The decay curves of $Ca_{0.8}Sr_{0.2}O:Ce^{3+}$, Eu^{2+} confirmed that there was an obvious energy transfer from Ce^{3+} to Eu^{2+} , which enhanced the quantum efficiency of $Ca_{0.8}Sr_{0.2}O:Eu^{2+}$. The thermal stability can also be enhanced by introducing Ce, which reduces the probability of non-radiative transitions at elevated temperatures, thereby minimizing electron loss. An NIR light emitting diode (LED) was fabricated by combining a blue LED chip with $Ca_{0.8}Sr_{0.2}O:Ce^{3+}$, Eu^{2+} , which was applied for the concentration detection of ethanol solutions.

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Introduction

In recent years, near-infrared (NIR) light has found widespread application in fields such as night vision, biological tissue penetration, solution detection, food quality detection, and plant growth. This is because NIR light possesses several advantageous properties: it has strong penetration capability, causes minimal damage to the human body, is invisible to the human eye, and overlaps with certain organic functional groups like C-H (2980 cm⁻¹) and C-O (1100 cm⁻¹).¹⁻⁹ As the core component for these applications, the NIR light source needs to meet the conditions of broadband emission, good stability, miniaturization and fast response.10 Traditional NIR light sources (such as the halogen tungsten lamp) can emit widerange NIR light; however, they have the disadvantages of high energy consumption, large volume and short life, which cannot meet the requirements for practical use.11,12 NIR LED is another NIR light source, which is made up of a combination of multiple NIR LEDs of different wavelengths arranged in an array to meet the demand of a broadband NIR light source. Although this NIR

Currently, NIR phosphors can be obtained by doping with the following ions: trivalent rare-earth ions (Pr³⁺, Yb³⁺, Nd³⁺), transition metal ions (Mn⁴⁺, Ni²⁺, Cr³⁺), and divalent rare-earth ions (Eu²⁺). Because electronic transitions inside the f-f electronic configuration of trivalent rare earth ions is forbidden and has a weak absorption capacity, such materials exhibit low luminous efficiency and a narrow emission bandwidth, which cannot meet the requirements of efficient wide-band NIR light sources, limiting their applications. Mn⁴⁺ ions tend to occupy a strong crystal field environment. They usually exhibit narrow band emissions, and the spectrum is not easy to regulate, thus lacking the requirements for a broadband emission NIR light source. Although the emission wavelength of Ni²⁺ ions is greater than 1000 nm, their luminescence efficiency is low with a relatively poor thermal stability, and it cannot be effectively applied as an NIR light source. Cr3+ exhibits a broadband NIR emission at ${}^{4}T_{2} \rightarrow {}^{4}A_{2}$ level transition. This transition is outside the 3d electron layer, and hence, it is vulnerable to the influence of

light source exhibits a long life and high efficiency, its large volume and the emission of various spectra by different LEDs consistently affect the overall spectral type of the light source with changes in temperature and time. These deficiencies make NIR LEDs unable to meet practical requirements. Currently, NIR phosphor conversion LEDs are considered ideal NIR light sources to satisfy the practical demands of the above-mentioned fields owing to their small size, long life and fast response. The main preparation method of an NIR pc-LED involves coating the broadband NIR phosphor onto a mature blue LED; hence, it is crucial to explore efficient broadband NIR phosphors.

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intensity changes in the crystal field. Consequently, Cr³⁺ will produce different luminescence under different conditions of crystal field strength. However, the Cr3+-doped NIR phosphors will inevitably contain toxic and harmful Cr6+, which greatly limits their application in food safety testing and human body

monitoring. The luminescence of Eu²⁺ (a rare earth ion) originates from its 5d \rightarrow 4f transition, and because of the naked leakage of the 5d level on the electron layer surface, Eu²⁺ is also susceptible to the crystal field environment, indicating that the luminescence of Eu²⁺ can be easily regulated. Therefore, Eu²⁺doped NIR phosphors have become a research hotspot in the field of NIR phosphors. In this work, a broadband NIR phosphor Ca_{0.8}Sr_{0.2}O:Eu²⁺ was prepared. It was observed that the energy transfer between Ce3+-Eu2+ was realized, quantum efficiency of Ca_{0.8}S_{r0.2}O:Eu²⁺ was improved from 36.7% to 45.8%, and the thermal stability was improved from 27.4% to 40.8%. The prepared NIR pc-LEDs were explored for the concentration detection of ethanol solutions. The results demonstrate that this study offers a novel approach for developing NIR phosphors.

2. **Experimental**

Paper

Two phosphors, namely, $Ca_{0.8}Sr_{0.2-x}O:xCe^{3+}$ and $Ca_{0.8-v}Sr_{0.2-}$ O:0.4% Ce³⁺, yEu²⁺ were prepared via a high-temperature solidstate method. High-purity (99.99%, Aladdin) CaCO₃, SrCO₃, Eu2O3, and CeO2 were used as raw materials, and they were precisely weighed in accordance with their stoichiometric ratios. The weighed samples were then placed in a clean agate mortar and ground thoroughly for 20-30 minutes. The ground powder was then transferred to either a corundum or graphite crucible and was placed in a tube furnace for high-temperature sintering. The temperature of the sintered samples was first raised from room temperature to 500 °C at 5 °C min⁻¹ for 30 minutes, then increased to 900 °C for 80 minutes, and finally elevated to 1200 °C for 60 minutes. The samples were allowed to cool to room temperature. Subsequently, they were ground again to form a homogeneous and fine powder for characterization and testing.

2.2. NIR pc-LED fabrication

Phosphor-converted LED (pc-LED): First, the phosphor powder, B glue, and A glue were weighed in a 5:4:1 ratio. Then, the weighed powder and glues were placed in a beaker for 40 minutes to ensure thorough mixing. Finally, the mixture was coated onto an LED chip with a suitable excitation wavelength and dried in an oven that was pre-set at 353 K for 2 hours to bond it to the chip.

Characterizations

The instrument used for the measurements was a Bruker AXS's D8 Advance X-ray diffractometer (Germany) with a Cu-Kα radiation source. The scan range was 10° to 80° (2θ) , operating at a precision of 0.0001°, with an applied voltage of 40 kV and a current of 40 mA. The structure was further optimized using the general structure analysis system (GSAS-II). Ultrahigh-resolution

surface micrographs of the nanoscale materials were acquired using an FEI Nova NanoSEM 450 scanning electron microscope, which also enabled the analysis of surface elemental composition (type and quantity). Surface analysis of the sample was conducted using an ESCALAB 250Xi X-ray photoelectron spectrometer, providing detailed information on the elemental composition, chemical states, molecular structures, and chemical bonding configurations of the phosphors. The analysis chamber operated at an ultimate vacuum of 5×10^{-1} mbar, with an energy scan range of 0 to 5000 eV and an energy resolution of 0.45 eV. The system featured independent analysis and injection chambers. A U4100 spectrometer with a wavelength range of 240 to 17 000 nm, a standard four-hole integrating sphere (with an opening rate of less than 7.8%), and a diameter of 60 mm was utilized, and it could accommodate a maximum sample size of 200×200 mm. Barium sulfate (BaSO₄) was used as the reference for transmission and absorption characterizations. Steady-state fluorescence emission spectra, excitation spectra, variable temperature spectra, and quantum efficiency were determined using a Fluorolog-3 fluorescence spectrometer (HORIBA Instruments, Inc.). The excitation source was a 450 W Xenon lamp, with a scanning speed of 1200 nm min^{-1} and an integration time of 0.1s.

Results and discussion

Phase information

Fig. 1(a) illustrates the layered arrangement of the CaO crystal structure. Fig. S1† and 1(b) show the XRD patterns of Ca_{0.8}- $Sr_{0.2-x}O:xCe^{3+}$ (x = 0.1-1.3%) and yEu^{2+} (y = 0%, 0.1%), and it can be seen that the synthesized samples matched with the CaO standard phase (PDF #001-1160) in the inorganic crystal structure database, indicating that the synthesized samples possessed a single phase. As we synthesized a sample with Sr²⁺, partially replacing Ca²⁺, the XRD pattern exhibited a tendency to be shifted in the direction of a small angle compared with the Standardized card. GSAS2 refinement software was employed to further investigate the effect of Ce3+-Eu2+ on the crystal structure of CSO. This refinement also aimed at determining whether the position of Eu²⁺ was changed. As exhibited in Fig. 1(c), for CSO:0.4% Ce³⁺ and CSO:0.4% Ce³⁺, 0.1% Eu²⁺, the profile factors $R_{\rm p}$, $R_{\rm wp}$, and χ^2 met the refinement requirements. This indicates that the crystal cell parameters obtained through refinement are reasonable, and this applies to other doping concentrations as well as $Ca_{0.8-x}Sr_{0.2}O:0.4\%$ Ce^{3+} , yEu^{2+} (y=0-0.12%). The crystal cell parameters obtained via refinement are depicted in Table 1. As depicted in Fig. 1(d), the lattice parameters a/b/c and volume V increase with the Eu²⁺ concentration. This indicates that Eu²⁺ ions still occupy the Ca²⁺ sites without changing their positions. Fig. 1(e) depicts the XPS spectra of CSO:0.4% Ce3+, 0.1% Eu2+; the binding energies corresponding to Ca-2p, Sr-3d, and O-1s can be well observed in the whole spectra. Owing to the small doping contents of Ce and Eu, they could only be observed in the fine spectrum, as depicted in the inset of Fig. 1(e). Additionally, the Eu-3d_{5/2} and the Ce³⁺ corresponding Ce-3d_{5/2} energy combinations were also observed. Fig. 1(f) presents the SEM image and elemental mapping of CSO:0.4% Ce³⁺, 0.1% Eu²⁺. The synthesized

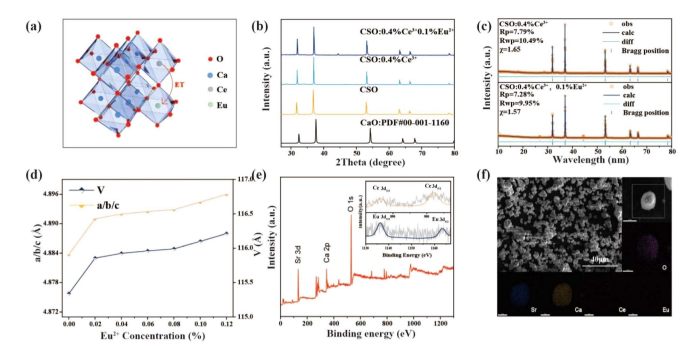


Fig. 1 (a) Crystal structure of CaO; (b) XRD patterns of $Ca_{0.8}Sr_{0.2}O$, $Ca_{0.8}Sr_{0.196}O(CSO)$:0.4% Ce^{3+} and CSO:0.4% Ce^{3+} , 0.1% Eu^{2+} ; (c) rietveld refinement of CSO:0.4% Ce^{3+} and CSO:0.4% Ce^{3+} , 0.1% Eu^{2+} ; (d) lattice parameters a/b/c and V with Eu^{2+} concentration; (e) XPS full survey spectrum of CSO:0.4% Ce^{3+} , 0.1% Eu^{2+} , inset: high-resolution XPS of Ce^{3+} and Eu^{2+} ; and (f) SEM and element mapping images of CSO:0.4% Ce^{3+} , 0.1% Eu^{2+} .

Table 1 Crystal cell parameters for $Ca_{0.8}Sr_{0.2}O:0.4\%$ Ce^{3+} , yEu^{2+} (y=0-0.12%) obtained *via* refinement

y(Eu ²⁺)	Cell parameters (ų)	Cell volume (ų)	$R_{\rm wp}$ (%), $R_{\rm p}$ (%), χ^2
0%	a/b/c = 4.87572	115.909	12.38, 8.5, 1.97
0.02%	a/b/c = 4.88299	116.428	14.59, 10.13, 2.08
0.04%	a/b/c = 4.88396	116.498	13.84, 9.8, 1.97
0.06%	a/b/c = 4.88446	116.533	12.56, 9.23, 1.76
0.08%	a/b/c = 4.88488	116.563	9.39, 6.79, 1.59
0.1%	a/b/c = 4.88638	116.671	9.95, 7.28, 1.57
0.12%	a/b/c = 4.88797	116.785	10.99, 9.8, 1.55

samples exhibited fairly uniform particle sizes of approximately 10 $\,\mu m$. Five elements, namely, Ca, Sr, O, Eu, and Ce were successfully identified.

3.2. Luminescence properties

Fig. 2(a) presents the emission spectrum of CSO:Ce³⁺ and the excitation spectrum of CSO:Eu²⁺. The spectral overlap between 500 nm and 700 nm suggested the potential energy transfer from Ce³⁺ to Eu²⁺. In order to further improve the CSO:Ce³⁺ performance by identifying the optimal Ce³⁺ concentration, the emission spectra of CSO:xCe³⁺ (x = 0.1-1.3%) were investigated under a 467 nm excitation, and the optimal Ce³⁺ concentration was obtained as x = 0.4% Fig. 2(b). Fig. 2(c) shows the emission spectra of CSO:0.4% Ce³⁺, yEu²⁺ (y = 0-0.12%) under an excitation of 467 nm, and the NIR emission peak belonging to Eu²⁺ appeared at y = 0.06%. Owing to concentration quenching, the NIR emission intensities first increased and then decreased as

the $\mathrm{Eu^{2^+}}$ concentration gradually increased, and the maximum value appeared at y=0.1%. Fig. 2(d) presents the decay curves of CSO:0.4% $\mathrm{Ce^{3^+}}$, $y\mathrm{Eu^{2^+}}$ (y=0–0.12%) at 581 nm, demonstrating the occurrence of energy transfer between $\mathrm{Ce^{3^+}}$ – $\mathrm{Eu^{2^+}}$, and all the decay curves could be well fitted using the biexponential function as follows: 16 -18</sup>

$$I(t) = I_0 + A_1 \exp\left(-\frac{t}{\tau_1}\right) + A_2 \exp\left(-\frac{t}{\tau_2}\right)$$
 (1)

where I(t) is the luminescence intensity, A_1 and A_2 are constants, τ is the average lifetime, and τ_1 and τ_2 are the fast decay and slow decay lifetimes, respectively. The average lifetime τ can be obtained using the following formula: 19,20

$$\tau = (A_1 \tau_1^2 + A_2 \tau_2^2) / (A_1 \tau_1 + A_2 \tau_2) \tag{2}$$

The calculations showed that the average lifetimes of Ce^{3+} ions were 33.48 ns, 28.46 ns, 20.53 ns, 13.73 ns, 10.59 ns, 5.01 ns and 4.68 ns for y=0%, 0.02%, 0.04%, 0.06%, 0.08%, 0.10%, and 0.12%, respectively. Clearly, the emission peak attributed to Ce^{3+} decreased monotonically with increasing Eu^{2+} concentration, proving the occurrence of energy transfer between Ce^{3+} Eu^{2+} . Energy transfer efficiency is an important parameter, which can be calculated by the decrease in emission intensity using the following formula:²¹

$$\eta_{\rm T} = 1 - I_{\rm s}/I_{\rm s0} \tag{3}$$

where I_{s0} and I_{s} represent the luminescence intensity of Ce³⁺ ions without and with Eu²⁺ ions, respectively, and η_{T} is the value

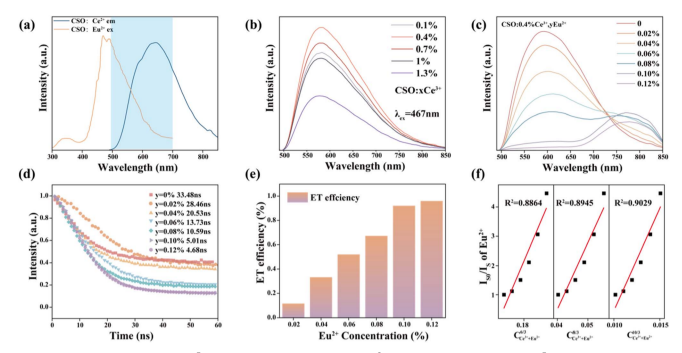


Fig. 2 (a) Emission spectrum of CSO:Ce³⁺ and excitation spectrum of CSO:Eu²⁺; (b) emission spectra of CSO:Ce³⁺ (x=0.1-1.3%); (c) emission spectra of CSO:0.4% Ce³⁺, yEu^{2+} (y=0-0.12%); (d) decay curves of CSO:0.4% Ce³⁺, yEu^{2+} (y=0-0.12%); (e) energy transfer efficiency of CSO:0.4% Ce³⁺, yEu^{2+} (y=0-0.12%) ($\lambda_{ex}=467$ nm); and (f) dependence of (l_{s0}/l_s) of Ce³⁺ on $C_{Ce+Eu}^{6/3}$, $C_{Ce+Eu}^{8/3}$ and $C_{Ce+Eu}^{10/3}$.

of energy transfer efficiency. The calculated results are depicted in Fig. 2(e). The energy transfer efficiency reached a maximum of 96.6% at x=0.12%.

A concentration quenching phenomenon can arise from either electric multipolar interactions or exchange interactions. When the critical distance exceeds 0.5 nm, the electric multipolar interactions are likely to induce concentration quenching; conversely, at shorter distances, the exchange interactions tend to dominate.²² As reported by Blaszse, the critical transfer distance (R_c) can be expressed as:²³

$$R_{\rm c} = 2[3V/4\pi X_{\rm c}N]^{1/3} \tag{4}$$

where N represents the number of host cations in the unit cell, V represents the volume per unit cell, and $X_{\rm c}$ represents the total critical concentration of ${\rm Ce^{3^+}}$ and ${\rm Eu^{2^+}}$ ions. For CSO substrate, $V=0.115909~{\rm nm^3}$, N=14, $X_{\rm c}=0.005$; therefore, $R_{\rm c}$ is approximately 1.468 nm, which is significantly larger than 0.5 nm, indicating that the electric multipolar interaction caused a nonradiative concentration quenching of ${\rm Ce^{3^+}-Eu^{2^+}}$ ions. For electrical multipolar interactions, it may include the dipoledipole (d–d), the dipole–quadrupole (d–q), and the quadrupole–quadrupole (q–q) interactions. According to Dexter's energy transfer theory, the multipolar interaction 24 , and the expressed using the linear relationship in the following formula:

$$\frac{\eta_0}{\eta_S} \propto C_{S+A}^{n/3} \tag{5}$$

where η_0 and η_S are the energy transfer efficiency of Ce³⁺ ions in the samples without and with Eu²⁺ ions, respectively. The relative luminescence intensity ratio (I_s/I_{s0}) can approximately

replace the ratio of η_0/η_s . $C_{\rm S+A}$ is the total concentration of ${\rm Ce}^{3+}$ and ${\rm Eu}^{2+}$ ions in the sample, and n=6, 8, or 10, indicating a d-d interaction, d-q interaction or q-q interaction, respectively. Fig. 2(f) depicts the linear relationship between $I_s/I_{\rm s0}$ and $C^{n/}$ $^4_{\rm S+A}$. An optimal linear behavior was obtained at n=10, indicating that q-q interactions determined the mechanism of energy transfer between ${\rm Ce}^{3+}$ and ${\rm Eu}^{2+}$ ions.

Fig. 3(a) displays the thermal spectra of CSO:0.4% Ce^{3+} , 0.1% Eu^{2+} . The emission intensity reached 40.8% (398 K) of that at room temperature (298 K), which is 13.4% higher than that of CSO:0.4% Ce^{3+} . To explore why thermal stability increased, Fig. 3(b) demonstrates the pyrothermal spectra of two samples before and after Ce^{3+} doping. Obviously, the pyrothermal curve intensity for the sample doped only with Eu^{2+} is nearly zero, indicating no trap. Conversely, after introducing isovalent Ce^{3+} , a peak appears in the pyroheat curve around 343 K. This suggests that blending Ce^{3+} – Eu^{2+} has created a shallow trap. The reason for occurrence of the trap was that the introduction of Ce^{3+} destroyed the original charge balance of the matrix. In order to maintain the charge balance, two Ce^{3+} replaced three Sr^{2+} to creating one doubly negatively charged Sr vacancy (V'_{Sr}) and two positively charged defects $[Ce^{3+}]_{Sr}^{-}$. The substitution process was as follows.

$$3Sr^{2+} + 2Ce^{3+} \rightarrow V_{Sr}^{"} + 2[Ce^{3+}]_{Sr}^{\cdot}$$
 (6)

The schematic of Ce^{3+} – Eu^{2+} energy transfer and thermal stability enhancement is given in Fig. 3(c). The electrons of Ce^{3+} jumped from the 4f ground state to 5d energy level after getting excited by a 467 nm light. A part of the electrons returned to the 4f ground state *via* radiative leaps and produced a broadband

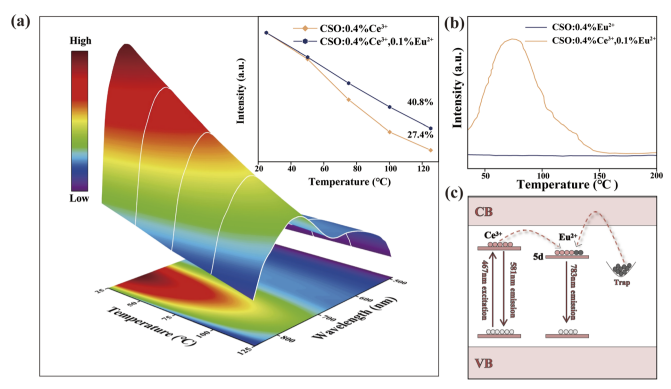


Fig. 3 (a) Temperature-dependent PL spectra of CSO:0.4% Ce^{3+} , 0.1% Eu^{2+} , inset: intensities of CSO:0.4% Ce^{3+} and CSO:0.4% Ce^{3+} , 0.1% Eu^{2+} at different temperatures; (b) TL curves of CSO:0.1% Eu^{2+} and CSO:0.4% Ce^{3+} , 0.1% Eu^{2+} ; and (c) mechanism diagram showing Ce^{3+} – Eu^{2+} energy transfer and thermal stability enhancement.

yellow luminescence, and another part of the electrons got transferred to the 5d energy level of Eu2+ via energy transfer and returned to the 4f ground state via radiative leaps, producing a broadband NIR emission. In addition, a small fraction of electrons was trapped and stored therein. In luminescent materials, nonradiative transitions are mainly caused by factors such as lattice vibrations and defects. When the temperature increases, the lattice vibration intensifies, which makes it easier for the electrons to return from the excited state to the ground state, thus increasing the probability of non-radiative transitions. In addition, defects in the material can have a significant effect on the non-radiative transitions. The presence of defects disrupts the periodic potential field of the lattice, creating localized potential wells, in which the electrons undergo nonradiative leaps. In this study, we observed that when the temperature was increased, the electrons stored in the traps jumped out of the traps owing to thermal excitation and moved through the conduction band to the 5d energy level of Eu²⁺. This process effectively replaced a part of the electron loss due to the increased probability of nonradiative leaps as a result of temperature elevation, resulting in an improvement in the thermal stability of the sample. In addition, we experimentally measured the quantum efficiency of the system to be 45.8% (as shown in Fig. S2†). This result also reflected that the material was thermally stable to a certain extent, and it can effectively suppress the effect of nonradiative transitions on the luminescence performance, demonstrating more intuitively the effects

of lattice vibrations and defect changes on nonradiative transitions.

4. Application

Organic solutions contain a large number of chemical bonds such as C-H, C-O and O-H; these chemical bonds can absorb near infrared light of specific wavelengths, and accordingly, the concentration of the organic solvents can be detected using an NIR light. An NIR pc-LED was prepared by combining a blue LED with CSO:0.4% Ce³⁺, 0.1% Eu²⁺ for concentration detection of ethanol (C2H5OH) solutions. Fig. 4(a) exhibits the experimental design for solution concentration detection. The NIR pc-LED emitted NIR light, which was passed through cuvettes with different concentrations of ethanol solution, and was finally received by the detector and presented on a computer in the form of a spectrum. Fig. 4(b) displays the experimental device. Fig. 4(c) exhibits the transmission spectra of the NIR pc-LED after passing through 0%, 20%, 40%, 60% and 80% ethanol solutions. It can be seen that as the concentration of the ethanol solution gradually increased, the absorption also gradually increased but the luminescence intensities gradually decreased. The intensity of the NIR light after passing through the ethanol solution and the concentration of the components in the sample complied with the Lambert-Beer law:

$$A = \log \frac{I_0}{I} = \varepsilon bc, \tag{7}$$

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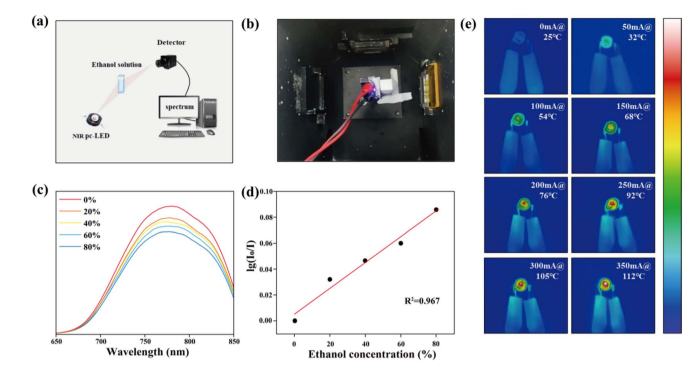


Fig. 4 (a) Schematic of the experimental design for solution concentration detection; (b) experimental device; (c) transmission spectra of the NIR pc-LED after passing through different concentrations of ethanol; (d) relationship between absorbance and concentration of ethanol solution; and (e) thermal images of the NIR pc-LED at different working currents ranging between 0–350 mA.

where A is the absorbance, I_0 is the initial intensity, I is the intensity after passing through different concentrations of ethanol solutions, ε denotes the molar absorptivity of the analyte, c is the concentration of the fraction to be tested, and b is the effective path length. The dependence of the luminescence intensity on the ethanol concentration was calculated. As illustrated in Fig. 4(d), the absorbance and ethanol concentration showed a good linear correlation after the fitting, and the fit reached $R^2 = 0.967$, indicating that the phosphor could offer a good application prospect in the field of concentration detection of solutions. The NIR pc-LED produced different amount of heat under different working currents, and thus, the temperature of the NIR pc-LED increased. Fig. 4(e) exhibits the pyrometer image of the NIR pc-LED under different working currents (0-350 mA). It can be seen that the temperature gradually increased with increasing working current, and when the working current reached 350 mA, the temperature reached 385 K.

5. Conclusions

In conclusion, the novel NIR phosphor $Ca_{0.8}Sr_{0.2}O:Eu^{2+}$ was synthesized via a solid-state method. When Ce^{3+} ions were introduced, the luminescence performance of $Ca_{0.8}Sr_{0.2}O:Ce^{3+}$, Eu^{2+} was obviously improved. Importantly, the quantum efficiency of $Ca_{0.8}Sr_{0.2}O:0.1\%$ Eu^{2+} increased from 36.7% to 45.8%, and the thermal stability at 398 K increased from 27.4% to 40.8%. The decay curves confirmed that the reason for this result was the occurrence of energy transfer between $Ce^{3+}-Eu^{2+}$. The NIR pc-LED prepared by combining $Ca_{0.8}Sr_{0.2}O:0.4\%$ Ce^{3+} ,

0.1% Eu²⁺ with a blue LED chip verified its promising application in the concentration detection of organic solutions.

Data availability

All relevant data are within the manuscript and its ESI.†

Conflicts of interest

The authors declare that they have no conflicts of interest.

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