Down-Conversion Polymer Composite Coatings with Multipeak Absorption and Emission

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Abstract: Spectral adjustment is an effective method to increase light conversion efficiency of solar cells and to promote the growth of plants. Down-converter (DC) materials are considered to be one of the most effective methods of spectral modification. The focus of this work was to expand the spectral response range of down-conversion layers to achieve multipeak absorption and emission. Sr₂CaMoO₆:Sm,Na and YVO₄:Bi,Eu, which have different excitation peaks in the UV-blue region and varied emission peaks in visible light regions, were prepared in this work. Sr₂CaMoO₆:Sm,Na can effectively produce red light at 648 nm upon excitation at 408 nm, while YVO₄:Bi,Eu can produce red light at 618 nm upon excitation at 365 nm. Polymeric luminescent coatings with one single kind of phosphor were prepared separately before the two phosphors were mixed together in uniform polymer coatings. The two phosphors were also assembled in bilayer coatings with different concentrations. The results showed that high transmittances over 90% were achieved for the two composite coatings with the thickness of 20 and 30 μm. The increase in particle loadings from 1% to 4% slightly decreased coating transmittance but increased luminescence intensity. The increase in the ratio of Sr₂CaMoO₆:Sm,Na and YVO₄:Bi,Eu from 5/1 to 10/1 resulted in high transmittance of the DC coatings, independent of total filler loadings (3‰ and 4‰) and coating thickness. The relative intensities of emission peaks can be adjusted conveniently by changing filler ratios. In addition, the transmittance and luminescent intensities of the coatings where the two phosphors were assembled in two layers were close to the uniform coatings, suggesting the negligible effect of UV light irradiation order. This work proved that the prepared coatings presented multipeak absorption and emission upon UV light excitation. These coatings can be expected to be applied in fields such as solar cells and agriculture greenhouses.

Keywords: Sr₂CaMoO₆:Sm,Na; YVO₄:Bi,Eu; polymeric luminescent coatings; down-conversion

1. Introduction

Spectral modification has been paid great attention in recent years because it has broad application prospects in many fields, including solar cells and agricultural greenhouses [1–4]. Taking solar cells as an example, spectral mismatch between uneven distribution of solar spectrum and spectral response of solar cells results in the limited conversion efficiency of solar cells [5,6]. In agriculture, shortwave photons that are not useful for plants can be converted into suitable light to promote their growth [7].

Down-conversion (DC) luminescent materials have become the focus of research as one of the spectral modification materials. [8–11]. Rare earth metals that have special electron layers and a rich number of energy level layers have been widely used in DC
luminescent materials to improve solar energy efficiency or promote plant growth. [12–16]. It is worth emphasizing that the rare earth ions used in down-conversion materials have no obvious negative impact on the environment, and different rare earth ions can be selected according to different spectral control requirements. Among them, rare earth organic complexes are composed of lanthanide ions and organic ligands exhibit great luminescence properties and have high-efficiency absorption in ultraviolet (UV) region [17,18]. However, the photochemical stability of lanthanide complexes limits their application. The high photodegradation rate upon UV exposure is a drawback that seriously affects their luminescence performance [19–21]. Inorganic phosphors as other potential DC materials are expected to be applied in many fields. Sulfides [15], nitride [22,23], nitrogen oxides [24,25] of rare earth metals are commonly used as red phosphors. Most of them are stable in physical and chemical properties, which is a big advantage over rare earth organic complexes. However, the narrow absorption band and low absorption coefficient limit inorganic phosphors’ further applications [26,27].

It is important to increase emission intensity and luminescent efficiency of inorganic phosphors [28]. Doping with different rare-earth ions can bring various luminescent effects [29]. The type and concentration of doping ions can be controlled to improve luminescence performance [30,31]. When rare-earth ion concentration in inorganic phosphors is higher than a critical valve, a quenching effect occurs and reduces the luminescent ability. In addition, defects in phosphor materials usually increase nonradiative transition channel and reduce quantum efficiency [32]. The reduction of defects in materials and nonradiative transition channels is a necessary approach to improve luminescent properties of phosphors [33,34]. Surface defects [35,36], atomic vacancy defects [37] and charge defects [38] usually exist in phosphors. Among them, charge defects often occur when trivalent rare earth ions are doped into the lattice to replace divalent metal ions. Such charge imbalance easily causes charge defects inside crystals. Several solutions are utilized to deal with this issue. For example, charge defects can be compensated in self-charge compensation by adding vacancies inside the lattice [39]. Another method is to replace divalent metal ions in crystal lattice with low-priced alkali metal ions (Li+, Na+, K+) [40,41]. The positive and negative charges in the lattice are mutually neutralized.

In addition to the increase in fluorescence intensity, the combination of two phosphors having different narrow absorption bands can be an effective method to achieve multiplex absorption and multiplex emission [42]. Molybdate and vanadate phosphors are candidate materials because of their different spectral characteristics and simple reaction conditions. Molybdate phosphors have low toxicity, low environmental pollution, good chemical stability, and good luminescent efficiency, which are potential candidates for red phosphors [43]. The MoO$_6^{6-}$ groups in molybdate phosphors can be excited by ultraviolet light in the range of 300–425 nm. The excitation range can be further adjusted to blue light band via the addition of Sm$^{3+}$ [44]. Vanadate phosphors are another functional inorganic material. Rare earth vanadate has high stability, good crystallinity and an effective absorption band in the ultraviolet region. Short-wavelength light energy can be converted into visible light by radiation due to the V–O charge transfer of VO$_4^{3-}$. Excellent fluorescence properties can be achieved when rare earth ions occupy Y$^{3+}$ sites in YVO$_4$ [45].

To date, most reports focus on DC materials with one single organic or inorganic phosphors [46]. When two phosphors are used together to achieve multiplex absorption and emission, the effects of total loading content, weight ratio, coating thickness on transparency, and luminescence should be concerned. More importantly, whether the distribution of two phosphors in coatings affects DC properties is not reported.

In this work, we prepared Sr$_2$CaMoO$_6$:Sm,Na and YVO$_4$:Bi,Eu that have different excitation peaks in UV-blue regions and various emission peaks in visible light regions before they were used together as phosphors in transparent luminescent polymer coatings to broaden the absorption of incident light and achieve an excellent emission spectrum. First, the luminescent properties of Sr$_2$CaMoO$_6$:Sm,Na red molybdate phosphors were improved by charge compensation. How the structures and their associated properties
varied with both amounts of Sm$^{3+}$ and Na$^+$ ions were then investigated. The effect of Na$^+$ ions was studied subsequently at a constant Sm$^{3+}$ ratio. The luminescent properties of YVO$_4$:Bi,Eu red phosphors were improved next by changing the concentration of rare-earth ions. The phosphor particles were further refined to improve their dispersion in a polymer matrix and to maintain transparency. Subsequently, luminescent polymeric coatings with one single kind of phosphors were prepared separately before the two phosphors were combined in uniform polymer coatings. For the ultraviolet light excitation sequence was concerned, the two phosphors were assembled in bilayer coatings with different concentrations. The transparency and luminescent spectra were investigated in detail.

2. Experimental

2.1. Materials and Synthesis of Sr$_2$CaMoO$_6$:Sm,Na and YVO$_4$:Bi,Eu

All the chemicals were commercially available. SrCO$_3$ (99%), CaCO$_3$ (99%), MoO$_3$ (99.9%), Sm$_2$O$_3$ (99.9%) and Na$_2$CO$_3$ (A.R. grade) were obtained from Sinopharm Chemical Reagent Co., Ltd. Y$_2$O$_3$ (99%), NH$_4$VO$_3$ (99%), Eu$_2$O$_3$ (99.9%) and Bi$_2$O$_3$ (99.9%) were all purchased from Aladdin Reagent Co., Ltd. Poly (Shanghai, China) (ethylene terephthalateco-1,4-cyclohexylenedimethylene terephthalate) (PETG) was provided by Eastman Kodak Chemical Company (Kingsport, TN, USA). 1,1,2,2-tetrachlor-ethane (C$_2$H$_2$Cl$_4$) was purchased from Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China).

Sr$_2$CaMoO$_6$:Sm,Na phosphors were synthesized by a high-temperature solid-state method with a tiny modification [47]. According to certain molar ratios, SrCO$_3$, CaCO$_3$, MoO$_3$, Sm$_2$O$_3$ and Na$_2$CO$_3$ were ground in a mortar for 40 min and heated in a muffle furnace at 600 $^\circ$C for 3 h. The calcined powder was ground in a mortar for another 30 min and further treated at 900 $^\circ$C for 3 h. Finally, the powder was placed in a high temperature electric furnace and heat-treated at 1200 $^\circ$C for 12 h. The name and the corresponding molar ratios of the examined samples are listed in Table 1.

Table 1. The names and corresponding molar ratios of raw materials for the examined Sr$_2$CaMoO$_6$:Sm,Na phosphors samples.

| Name    | SrCO$_3$ | CaCO$_3$ | MoO$_3$ | Sm$_2$O$_3$ | Na$_2$CO$_3$ |
|---------|----------|----------|----------|-------------|--------------|
| Sm20Na00| 1.980    | 1        | 1        | 0.0100      | 0.0000       |
| Sm10Na10| 1.980    | 1        | 1        | 0.0050      | 0.0050       |
| Sm15Na15| 1.970    | 1        | 1        | 0.0075      | 0.0075       |
| Sm20Na20| 1.960    | 1        | 1        | 0.0100      | 0.0100       |
| Sm25Na25| 1.950    | 1        | 1        | 0.0125      | 0.0125       |
| Sm30Na30| 1.940    | 1        | 1        | 0.0150      | 0.0150       |
| Sm35Na00| 1.975    | 1        | 1        | 0.0125      | 0.0000       |
| Sm25Na15| 1.960    | 1        | 1        | 0.0125      | 0.0075       |
| Sm25Na20| 1.955    | 1        | 1        | 0.0125      | 0.0100       |
| Sm25Na30| 1.945    | 1        | 1        | 0.0125      | 0.0150       |
| Sm25Na35| 1.940    | 1        | 1        | 0.0125      | 0.0175       |

YVO$_4$:0.04Bi$^{3+}$, Eu$^{3+}$ phosphors were synthesized via a solid-state reaction method. According to certain molar ratios, Y$_2$O$_3$, NH$_4$VO$_3$, Eu$_2$O$_3$, Bi$_2$O$_3$ were mixed and ground in an agate mortar with a pestle. Subsequently, the well-ground mixture was calcined at 800 $^\circ$C for 4 h in an air atmosphere. The name and the corresponding molar ratios of the examined samples are listed in Table 2.

The obtained nanophosphors were ground for another 48 h and used for a further measurement.
Table 2. The names and corresponding molar ratios of raw materials for the examined YVO₄:0.04Bi³⁺, Eu²⁺ samples.

| Name  | Y₂O₃   | NH₄VO₃ | Bi₂O₃  | Eu₂O₃  |
|-------|--------|--------|--------|--------|
| Eu01  | 0.475  | 1      | 0.02   | 0.005  |
| Eu02  | 0.470  | 1      | 0.02   | 0.010  |
| Eu03  | 0.465  | 1      | 0.02   | 0.015  |
| Eu04  | 0.460  | 1      | 0.02   | 0.020  |
| Eu05  | 0.455  | 1      | 0.02   | 0.025  |

2.2. Preparations of the Polymeric Luminescent Coatings

The luminescent polymeric coatings with single phosphors were prepared as follows. The phosphor was dispersed ultrasonically in 100 mL C₂H₂Cl₄ before 20 g of PETG were added. The mixture was stirred at 80 °C until PETG was completely dissolved. The mixed solutions containing different phosphor concentrations (1–4‰) were obtained. Then, a spray coating method was used for preparing the coatings on quartz glass. The thickness was controlled to be 20 or 30 μm, respectively.

The single-layer luminescent polymeric coatings that were directly mixed with two phosphors were prepared as well. The weight ratios of YVO₄:Bi,Eu and Sr₂CaMoO₄:Sm,Na were 1:5 and 1:10, respectively. The total powder contents were 3‰ or 4‰. The following preparation of coatings was the same as mentioned above.

Double-layer coatings were prepared, and the total powder contents were also 3‰ or 4‰. The coating thickness was 30 μm. The two phosphors were distributed in different layers. The names of the polymeric luminescent coatings and corresponding parameters are listed in Tables 3–5.

Table 3. The names of the polymeric luminescent coatings with single phosphors.

| Name a | Sm25Na25 Weight Ratio | Thickness b (μm) | Name a | Eu04 Weight Ratio | Thickness b (μm) |
|--------|-----------------------|------------------|--------|-------------------|------------------|
| M-1T20 | 1‰                   | 20               | V-1T20 | 1‰               | 20               |
| M-2T20 | 2‰                   | 20               | V-2T20 | 2‰               | 20               |
| M-3T20 | 3‰                   | 20               | V-3T20 | 3‰               | 20               |
| M-4T20 | 4‰                   | 20               | V-4T20 | 4‰               | 20               |
| M-1T30 | 1‰                   | 30               | V-1T30 | 1‰               | 30               |
| M-2T30 | 2‰                   | 30               | V-2T30 | 2‰               | 30               |
| M-3T30 | 3‰                   | 30               | V-3T30 | 3‰               | 30               |
| M-4T30 | 4‰                   | 30               | V-4T30 | 4‰               | 30               |

a: M and V stand for molybdate and vanadate phosphors, respectively. T stands for thickness in μm. b: The uncertainty of coating thickness was ±1 μm.

2.3. Characterizations and Calculation

An X-ray diffractometer (ARLXTRAX-3KW, American Thermoelectric Company, Pennsylvania, Massachusetts, America) was used to characterize the phase, crystal structure, and unit cell parameters of the prepared samples at a scan rate of 10° min⁻¹ and a scan range of 10–80°. A scanning electron microscope (JSM-IT200, Tokyo, Japan) was used to observe the surface topography of the sample. Agilent Cary 5000 UV-Vis-NIR spectrophotometer was used for spectral testing of the prepared samples. Reflectance test conditions were set to a slit width of 2 nm, a scan speed of medium speed and a test wavelength of 200–1000 nm. The fluorescence performance was tested at a certain excita-
tion wavelength by an FL3-221 fluorescence spectrophotometer from HORIBA JobinYvon, France. The particle size distribution of phosphors was obtained by a laser diffraction particle size analyzer (Mastersizer 2000, Malvern Instruments Ltd., Worcestershire, UK).

Table 4. The name of the polymeric luminescent coatings that were directly mixed with two phosphors.

| Name a | Ratio (Eu04:Sm25Na25) | Total Powder Weight Ratio | Thickness b (µm) |
|--------|----------------------|--------------------------|------------------|
| Z3R15T20 | 1:5 | 3‰ | 20 |
| Z3R15T30 | 1:5 | 3‰ | 30 |
| Z3R110T20 | 1:10 | 3‰ | 20 |
| Z3R110T30 | 1:10 | 3‰ | 30 |
| Z4R15T20 | 1:5 | 4‰ | 20 |
| Z4R15T30 | 1:5 | 4‰ | 30 |
| Z4R110T20 | 1:10 | 4‰ | 20 |
| Z4R110T30 | 1:10 | 4‰ | 30 |

Note: Z means that vanadate and molybdate phosphors were mixed in polymer coatings. R and T stand for their weight ratio and coating thickness, respectively. b: The uncertainty of coating thickness was ±1 µm.

Table 5. The name of the double-layer polymeric luminescent coatings with different phosphor weight ratios.

| Name a | Total Powder Weight Ratio | Ratio (Eu04:Sm25Na25) | Upper Layer | Lower Layer |
|--------|--------------------------|----------------------|-------------|-------------|
| Bi3R15SE | 3‰ | 1:5 | Sm25Na25 | Eu04 |
| Bi3R110SE | 3‰ | 1:10 | Sm25Na25 | Eu04 |
| Bi4R15SE | 4‰ | 1:5 | Sm25Na25 | Eu04 |
| Bi4R110SE | 4‰ | 1:10 | Sm25Na25 | Eu04 |
| Bi3R15ES | 3‰ | 1:5 | Eu04 | Sm25Na25 |
| Bi3R110ES | 3‰ | 1:10 | Eu04 | Sm25Na25 |
| Bi4R15ES | 4‰ | 1:5 | Eu04 | Sm25Na25 |
| Bi4R110ES | 4‰ | 1:10 | Eu04 | Sm25Na25 |

Note: a: Bi means that vanadate and molybdate phosphors were mixed separately in bilayered polymer coatings. R stands for their weight ratio. The whole coating thickness was 30 ± 1 µm.

The calculation of the electronic structures of YVO₄:Bi and Sr₂CaMoO₆:Sm₃Na was performed with the density functional theory (DFT) using the CASTEP code, in which the electron–ion interactions were described by pseudopotential method and electronic wave functions are represented by means of a plane-wave basis set. The first step was to use crystallographic data to construct a YVO₄ unit cell, in which one Y ion is replaced by a Bi ion. Generalized gradient approximation (GGA) by the Perdew, Burke and Ernzerhof (PBE) formulation was chosen to optimize the crystal structure. The second step was to calculate the band structure and density of states of YVO₄ and YVO₄:Bi. The kinetic cutoff energy was 571 eV and Brillouin zone integration was represented using the K-point sampling scheme of 5 × 6 × 6 Monkhorst-Pack grid. Ultrasoft pseudopotentials were used...
to approximate the core electrons. The calculation method and setting parameters of the Sr$_2$CaMoO$_6$:Sm and Sr$_2$CaMoO$_6$:Sm,Na are the same as those of YVO$_4$:Bi.

3. Results and Discussion

3.1. Sr$_2$CaMoO$_6$:Sm,Na Phosphors

3.1.1. Effect of Doping Amounts of Sm$^{3+}$ and Na$^+$

The effect of Sm$^{3+}$ and Na$^+$ doping was studied first. The XRD patterns of the obtained phosphor samples of Sr$_2$CaMoO$_6$:Sm,Na and the JCPDS card No: 48-799 are shown in Figure 1a. Main diffraction peaks at 30.8° and 44.1° of the phosphors were observed, which match with the JCPDS standard card peaks of α-Sr$_2$CaMoO$_6$ with an orthogonal crystal system. Because the radius of the doped Sm$^{3+}$ and Na$^+$ ions are close to the radius of Sr$^{2+}$ ions, they can occupy the lattice sites of Sr$^{2+}$ at random. The diffraction peaks in Sm$_{10}$Na$_{10}$–Sm$_{30}$Na$_{30}$ doped with Na$^+$ were not changed compared with those in Sm$_{25}$Na$_{00}$. It can be argued that Sm$^{3+}$ and Na$^+$ may enter the matrix lattice of Sr$_2$CaMoO$_6$ effectively.

![Figure 1](image-url)  

Figure 1. (a) XRD patterns of the Sr$_2$CaMoO$_6$:Sm,Na (Sm$_{20}$Na$_{00}$–Sm$_{30}$Na$_{30}$). (b) Excitation spectra ($\lambda_{em}$ = 648 nm) of the Sr$_2$CaMoO$_6$:Sm,Na samples. (c) Fluorescence emission spectra of Sr$_2$CaMoO$_6$:Sm,Na samples ($\lambda_{ex}$ = 408 nm). (d) Fluorescence emission spectra of Sr$_{1.975}$CaMoO$_6$:0.025Sm$_x$Na$_x$ series samples. ($\lambda_{em}$ stands for the emission wavelength and $\lambda_{ex}$ stands for the excitation wavelength.).

The excitation spectra (Figure 1b) were composed of two parts in the UV-blue region. The wide peak at 350–450 nm originated from the electron transition of O$^{2-}$ → Mo$^{6+}$ charge transfer band in the MoO$_6^{6-}$ matrix. The other part of narrow-band excitation peaks was located at 450–500, 530, and 565 nm due to the characteristic transition of Sm$^{3+}$ from ground state $^6$H$_{5/2}$ to various high-energy excited states. The luminescence intensities at the highest excitation peak (408 nm) are 14 and 5.2 times higher than those of the phosphors without charge compensation ion Na$^+$ when the doping amounts of Sm$^{3+}$ and Na$^+$ were 2.5 mol% and 1 mol%, respectively. The luminescence intensities at the excitation peak of 465 nm are 5 and 1.8 times higher than those of the phosphors without charge compensation ion Na when the doping amounts of Sm$^{3+}$ and Na$^+$ were 2.5 mol% and 1 mol%, respectively. The fluorescence emission spectra of Sr$_2$CaMoO$_6$:Sm,Na samples are shown in Figure 1c. The emission spectra excited by 408 nm are mainly the characteristic
peaks of Sm$^{3+}$. The strong emission peak at 648 nm and the relatively weak emission peaks at 562, 600 and 712 nm were generated by $^4G_{5/2} \rightarrow ^6H_{9/2}$, $^4G_{5/2} \rightarrow ^6H_{5/2}$, $^4G_{5/2} \rightarrow ^6H_{7/2}$, and $^4G_{5/2} \rightarrow ^6H_{11/2}$ energy level transitions, respectively. The fluorescence intensity of Sm$^{25}$Na$^{25}$ reached the highest value, which is five times higher than that of the undoped sample. This result is also consistent with the excitation spectrum (Figure 1b).

To verify whether the low-cost alkali metal Na$^+$ had the charge compensation effect, the phosphors of Sr$_{1.975-x}$CaMoO$_6$:0.025Sm$_x$Na series were synthesized by only changing the doping amount of Na$^+$ (Table 1). The fluorescence properties of Sr$_{1.975-x}$CaMoO$_6$:0.025Sm$_x$Na samples prepared by high-temperature solid phase are shown in Figure 1d. The emission intensity was enhanced by increasing the Na$^+$ content of the samples, when they were excited by the light at 408 nm. By further increasing Na$^+$ concentration, the excessive charge caused by the compensator brought lattice defects due to charge imbalance in the lattice, decreasing the internal quantum efficiency. The maximum intensity, thus, was obtained until the doping amounts of Na$^+$ and Sm$^{3+}$ were equal in Sm$^{25}$Na$^{25}$.

3.1.2. Band Structure and Density of States

For the purpose of clarifying the role of Na$^+$ on the band structure in the phosphor host, we calculated the band structure and total density of states (DOS) of the Sm$^{25}$Na$^{00}$ and Sm$^{25}$Na$^{25}$. The calculated electronic band structures of the phosphors on the basis of DFT calculations are shown in Figure 2. It can be observed that the energy structures of the two samples are similar. The lowest energy of conduction bands (CBs) was located at S site of Brillouin zone and the highest point of valence bands (VBs) was located at G point. Hence, the phosphors were indirect bandgap materials. The gap between the lowest energy of the CBs and the highest energy of the VBs in Sm$^{25}$Na$^{25}$ was 2.063 eV, which is 0.018 eV smaller than that of the optical band gap of Sm$^{25}$Na$^{00}$. The reason for the small difference in optical band gap between the two samples is the low doping amount of Na$^+$. The DOS of the phosphors is shown in Figure 2. After doping with Na$^+$, the main peak in density of states in the Sm$^{25}$Na$^{25}$ were improved compared with the nondoped sample, suggesting that the free electrons were increased. The above calculation results also agree with the experimental results, i.e., the addition of Na$^+$ enhanced the luminescence properties of the phosphors.

![Figure 2](image-url)
3.2. YVO$_4$:Bi,Eu

3.2.1. Effect of Amounts of Eu$^{3+}$

The XRD patterns of the samples with different Eu$^{3+}$ doping concentrations and the JCPDS card No: 17-0341 are shown in Figure 3a. These samples are completely consistent with tetragonal YVO$_4$. The doping of Eu$^{3+}$ ions did not cause any significant changes in the crystal structure. The diffraction pattern reveals that Eu$^{3+}$ ions were substituted in the Y$^{3+}$ sites of YVO$_4$ host. In addition, impurity peaks were negligibly detected within the experimental range and no shift in peaks for Eu$^{3+}$ doped YVO$_4$ was found, indicating that the samples are mainly in the pure tetragonal phase.

![Figure 3. (a) XRD patterns of YVO$_4$:Bi,Eu. (b) Excitation ($\lambda_{em} = 618$ nm) spectra of YVO$_4$:Bi,Eu. (c) Emission ($\lambda_{ex} = 365$ nm) spectra of YVO$_4$:Bi,Eu.](image)

To obtain the sample with optimal luminescence properties, the excitation and emission spectra of samples were characterized. As shown in Figure 3b, a broad excitation band appeared in the range of 250–400 nm that corresponds to the charge transfer from excited ligands to the central vanadium atom inside VO$_4^{3-}$ groups. A weak peak at 395 nm originated from the $^7F_0-^5L_6$ transitions within the 4f$^6$ configuration of the Eu$^{3+}$ ions. The intensity reached the highest when the Eu$^{3+}$ concentration was 4 mol%. In the emission spectra (Figure 3c), the weak emission at 593 nm and strong emission at 618 nm correspond to the $^5D_1-^7F_1$ and $^5D_0-^7F_j$ ($j = 0–4$) transition of Eu$^{3+}$, respectively. The emission intensity gradually increased when the concentration of Eu$^{3+}$ varied from 1 to 4 mol%. The intensity reached the highest level at 4 mol%, and then the intensity decreased with the further increased concentration due to the concentration quenching effect. Therefore, the sample Eu04 was used for further study.

3.2.2. Band Structure and Density of States

To clarify the role of Bi on the band structure in YVO$_4$:Bi,Eu host, band structure and partial density of states (PDOS) of atoms were calculated to explain chemical bonding properties and origin of CT transitions of YVO$_4$:Bi,Eu. The DFT method was also used to calculate the influence of the replacement of Y with Bi in the YVO$_4$:Bi,Eu host. A unit cell with 20 atoms was used, and the cutoff energy was 571 eV. One Y atom is substituted by Bi. Due to the large and expensive computation, the cases with the Bi percentage were not simulated. Figure 4 shows the calculated band structure along the high symmetry points.
of the first Brillouin zone for YVO$_4$:Bi. The lowest energy of conduction bands (CBs) was localized at G point, while the highest point of the valence bands (VBs) was in P point. Hence, YVO$_4$:Bi is an indirect bandgap material, and the gap between the lowest energy of the CBs, and the highest energy of the VBs is about 2.846 eV. DFT is a ground state theory, but band gap belongs to the properties of excited states. Therefore, the difference from generalized gradient approximation (GGA) in DFT calculations generally underestimates the band gap of insulators and semiconductors.

![Energy Band Structure and Density of States](image)

**Figure 4.** The calculated energy band structure and density of states of YVO$_4$:Bi,Eu (a,b). Partial densities of states of YVO$_4$:Bi,Eu (c-f) near the Fermi energy level (the Fermi energy is the zero of the energy scale).

The partial density of states (PDOS) of YVO$_4$:Bi is illustrated in Figure 4 to explain the origin of the CBs and the highest energy of VBs. The conduction band in the range of 2.0–6.0 eV is dominated by V-3d, mixed with a small amount of Bi-6p and O-2p states. The valence band just below the Fermi level (0.0 eV) was mainly occupied by O-2p and V-3d, with some Bi-6s and Bi-6p states’ contributions. Therefore, the absorption in the UV region can be ascribed to the CT transitions from O-2p to V-3d and Bi-6p states, respectively. Combined with the optical band gap calculations, it can be argued that efficient absorption in the near UV region can exist even when the content of Bi is very low because of its occupation in the lowest conduction band of YVO$_4$:Bi host.
3.3. Influence of Particle Size

It is also challenging to achieve good dispersion of down-conversion materials in an appropriate polymer matrix that must exhibit high transmittance to prevent scattering and absorption. The phosphor samples Sm25Na25 and Eu04 were ground into fine powders subsequently (see experimental section), and the size effect on the fluorescence performance was studied.

Figure 5a,b shows the particle size distributions and scanning electron micrographs of Sm25Na25 phosphors. The initial size of the phosphor was about 30 μm (Figure 5a). As shown in Figure 5b, the particle size of the phosphor was reduced, and the distribution was between 2−10 μm. X-ray diffraction patterns of samples with different sizes are shown in Figure 5c. The position and intensity of the XRD diffraction peaks did not change significantly, indicating that the crystal structure of the phosphors was not destroyed after ball milling. As shown in the fluorescence emission spectra of the two samples (Figure 5d), the fluorescence intensity of the large phosphor is significantly higher than that of the small one. The crystal defects that were generated during the grinding process increased the nonradiative transition channel, thereby weakening the luminescence performance of the phosphor.

![Figure 5](image_url)

**Figure 5.** (a) The initial particle size distribution of Sm25Na25. (b) Particle size distribution of Sm25Na25 after grinding. (c) XRD spectra before and after grinding. (d) Emission spectra before and after grinding.

The particle size distributions and scanning electron micrographs of Eu04 are shown in Figure 6. It is evident that particles had irregular shapes and wide size distributions (Figure 6a). The particle size ranged from 1 to 14 μm. As shown in Figure 6b, particle size was reduced to the nanometer level, and the size distribution was mainly concentrated in 300–600 nm. The X-ray diffraction patterns and luminous performance of samples with different sizes are shown in Figure 6c,d. Similarly, the phosphors before and after ball milling had higher crystallinity, but the smaller size brought more crystal defects and reduced the fluorescence intensity.
3.4. Preparations and Properties of Luminescent Polymeric Coatings

The luminescent coatings can be obtained by dispersing luminescent materials in polymer coatings. Therefore, luminescent polymeric coatings with one single kind of phosphors were prepared. In order to broaden the absorption of the spectrum, the two phosphors were combined together in uniform polymer coatings. As far as the UV light excitation sequence was concerned, the two phosphors were assembled in bilayer coatings with different concentrations.

Figure 7a,b shows the transmittance of PETG coatings with different phosphor concentrations (1–4‰) and coating thickness (20 and 30 µm). High transmittance >92% can be observed in visible-infrared regions, which would hardly influence the light transmittance of the coatings. The coating transmittance was decreased as the coating thickness increased. The characteristic emission peaks were found at 562, 600, and 648 nm upon the excitation of 408 nm. Photoluminescence emission (PL) spectra also showed high intensities for coatings with high phosphor concentration and coating thickness, especially the main emission peak at 648 nm. The optical properties of YVO₄:Bi,Eu (1, 2, 3 and 4‰) in PETG coatings were examined as well (Figure 7c,d). Similar results were also obtained. The transmittance of the composite coatings was decreased with phosphor concentration and coating thickness, while the luminescent intensity was increased. The luminescent coatings showed the YVO₄:Bi,Eu characteristic emission peaks at 593, 615, 618, 650 and 698 nm upon the excitation at 365 nm.

The different excitation and emission wavelengths of the two phosphors enable their combination to broaden the absorption band of the UV light in the solar spectrum and achieve multipeak emission in the visible light band. Herein, the single-layer luminescent polymeric coatings that were directly mixed with two phosphors were prepared. The transmittances of coatings with two different phosphors are shown in Figure 8. High transmittance in the visible-infrared region can still be observed. The increase in coating thickness with the same phosphor concentration decreased the transmittance. The transmittance decreased as the phosphor content increased when the coating thickness was the same. Keeping the total phosphor content unchanged, the coating with more Sm25Na25 had higher transmittance than the coating with less Sm25Na25.
PL emission spectra of coatings are shown in Figure 9. The PL spectrum displayed the emissions of YVO$_4$:Bi,Eu and Sr$_2$CaMoO$_6$:Sm,Na. Upon excitation at 365 nm, the coatings showed the characteristic emission bands of Eu$^{3+}$ ions. The weak emission at 593 nm and the strong emission at 618 nm correspond to the $^5$D$_{0}$-$^7$F$_J$ and $^5$D$_{0}$-$^7$F$_J$ ($J = 0$–4) transition. The fluorescence intensity increased significantly as the powder content and coating thickness were increased. The emission at 650 nm excited by 365 nm is contributed by the two phosphors. Upon the excitation at 408 nm, the coatings showed the weak emission at 600 nm and the strong emission at 648 nm. Fluorescence intensity at 648 nm increased significantly with the increase in Sr$_2$CaMoO$_6$:Sm,Na content. The results proved that the coatings with two phosphors should be an ideal luminescence downshifting (LDS) layer.
Coatings showed the characteristic emission bands of Eu3+ ions. The weak emission at 593 nm and the strong emission at 618 nm correspond to the 5D1 – 7F1 and 5D0 – 7FJ (J = 0 – 4) transition. The fluorescence intensity increased significantly as the powder content and coating thickness were increased. The emission at 650 nm excited by 365 nm is contributed by the two phosphors. Upon the excitation at 408 nm, the coatings showed the weak emission at 600 nm and the strong emission at 648 nm. Fluorescence intensity at 648 nm increased significantly with the increase in Sr2CaMoO6:Sm,Na content. The results proved that the coatings with two phosphors should be an ideal luminescence downshifting (LDS) layer.

Figure 9. (a) Emission spectra of Z3R15T20 and Z3R15T30 upon the excitation at 365 and 408 nm, respectively. (b) Emission spectra of Z3R110T20 and Z3R110T30. (c) Emission spectra of Z4R15T20 and Z4R15T30. (d) Emission spectra of Z4R110T20 and Z4R110T30.

Since vanadate and molybdate are excited by 365 and 408 nm photons, respectively, the penetration depth of the ultraviolet light is different. Therefore, the sequence of vanadate and molybdate may affect the fluorescence intensity of the coatings. The two phosphors were assembled in bilayer coatings with different concentrations. As shown in Figure 10a,b, high transmittance of >92% was observed in visible-infrared regions from the transmittance spectrum. Compared with the single-layer coatings, the double-layer coatings had the same high transmittance. Figure 10c–f shows fluorescence emission spectra of the bilayer coatings. It is obvious that the coatings exhibited multiple emission peaks in the visible region. Similar to the single-layer coatings directly mixed with two phosphors, the double-layer coatings exhibited the characteristic emission peaks of different phosphors under excitation at 365 and 408 nm. The emission peak at 593 nm and 618 nm and the emission at 650 nm which were produced by the two powders together guaranteed multipeak emission. From the emission spectrum that the corresponding fluorescence intensity was higher when YVO4:Bi,Eu was in the upper layer. On the contrary, when Sr2CaMoO6:Sm,Na was in the upper layer, its corresponding emission intensity was increased. The difference in fluorescence performance controlled by layer sequence provides a way to adjust the coating’s DC performance.
Figure 10. (a−b) Transmission spectrum of the bilayer coatings. (c) Emission spectra of Bi3R15SE and Bi3R15ES upon the excitation at 365 and 408 nm, respectively. (d) Emission spectra of Bi3R110SE and Bi3R110ES. (e) Emission spectra of Bi4R15SE and Bi3R15ES. (f) Emission spectra of Bi4R110SE and Bi4R110ES.

4. Conclusions

In this paper, Sr2CaMoO6:Sm,Na and YVO4:Bi,Eu that have different excitation in the UV-blue region and varied emission peaks in the visible light regions were synthesized. Sr2CaMoO6:Sm,Na can effectively produce red light at 648 nm upon the excitation at 408 nm. The luminescent properties of Sr2CaMoO6:Sm,Na red molybdate phosphors were improved by charge compensation. Subsequently, by changing the concentration of rare-earth ions, the absorption capacity of YVO4:Bi, Eu red molybdate phosphor in the ultraviolet region and the luminous intensity in the visible light region were improved. YVO4:Bi,Eu can effectively produce red light at 618 nm upon the excitation at 365 nm. The DFT method was used to explain the chemical bonding properties and origin of CT transitions of YVO4:Bi,Eu. Next, the polymeric luminescent coatings with a single phosphor were prepared. The coating’s DC ability increased with the increase of phosphors concentration, and the transmittance was above 92%. In order to achieve the effects of multipeak absorption and wide spectrum emission, the two phosphors were combined together in uniform polymer coatings. The coatings exhibited the characteristic fluorescence peaks of Sr2CaMoO6:Sm,Na and YVO4:Bi,Eu upon excitation at 365 nm and 408 nm, respectively. The two phosphors were assembled in bilayer coatings with different concentrations, and the transparency and luminescent spectra were investigated in detail. High transmittances over 90% were achieved for the two composite coatings with the thickness of 20 and 30 μm. The increase in particle loadings from 1‰ to 4‰ slightly decreased film transmittance but
increased luminescence intensity, independent of total filler loadings (3% and 4%) and coating thickness. The relative intensities of emission peaks can be adjusted conveniently by changing filler ratios. Similar to the single-layer coatings directly mixed with two phosphors, the bilayer coatings have high transmittance and the characteristic emission peaks of different phosphors under excitation at 365 and 408 nm. The transmittance and luminescent intensities of the coatings where the two phosphors were assembled in two layers were close to the uniform coatings, suggesting the negligible effect of UV light absorption order. The results proved that the coatings with multipeak absorption and multipeak emission are expected to be applied in the fields of solar cells and agricultural greenhouses.

**Author Contributions:** Conceptualization, C.Z. and C.L.; methodology, J.F.; software, J.F.; validation, J.F.; formal analysis, J.F. and W.L.; investigation, W.Z.; resources, Y.N.; data curation, J.F.; writing—original draft preparation, J.F. and L.F.; writing—review and editing, C.Z.; visualization, J.F.; supervision, C.L.; project administration, C.L.; funding acquisition, C.L. All authors have read and agreed to the published version of the manuscript.

**Funding:** This research was funded by Natural Science Foundation of Jiangsu Province, No. BK20191364, No. BK20141459.

**Institutional Review Board Statement:** Not applicable.

**Informed Consent Statement:** Not applicable.

**Data Availability Statement:** The data presented in this study are available in this article.

**Acknowledgments:** The project was also funded by the Priority Academic Program Development of the Jiangsu Higher Education Institutions (PAPD) and Jiangsu Collaborative Innovation Center for Advanced Inorganic Function Composites.

**Conflicts of Interest:** The authors declare that they do not at present have and do not anticipate having any conflict of interests. All authors have the authority to submit this work for publication. The manuscript or its contents in some other form have not been published previously by any of the authors and/or is not under consulting for publication in another journal at any time of the submission.

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