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## 1. Introduction

In recent years, the spectroscopy and optical properties of chromium single-doped and chromium/rare-earth (RE) codoped have been extensively studied<sup>1-5</sup> due to their advantages.6,7 Chromium is a transition metal (TM) with many different valence states. $8-10$  In the glass networks, it often exists in the trivalent state  $Cr^{3+}, ^{1-5,8}$  which can emit radiation in the visible (VIS), near-infrared (NIR) regions under different excitation wavelengths.<sup>5,11,12</sup> Also, the <sup>4</sup>T<sub>1g</sub>(F)  $\rightarrow$  <sup>4</sup>A<sub>2g</sub>, <sup>4</sup>T<sub>2g</sub>(F)  $\rightarrow$  <sup>4</sup>A<sub>2</sub>  $\rightarrow$  <sup>4</sup>A<sub>2</sub><sup>2</sup>E  $\rightarrow$  <sup>4</sup>A<sub>2</sub><sup>4</sup>A<sub>2</sub> transitions of Cr<sup>3+</sup>  $\rm A_{2g}, {}^2T_{1g} \rightarrow {}^4A_{2g}, {}^2T_{2g} \rightarrow {}^4A_{2g}$  and  ${}^2E \rightarrow {}^4A_{2g}$  transitions of  $\rm Cr^{3+}$ can be combined with  $Yb^{3+}$  to generate the VIS, NIR emission spectra.<sup>5,13,14</sup> In 2001, H. U. Güdel et al.<sup>15</sup> confirmed that  $Cr^{3+}$ , in association with  $Yb^{3+}$ , creates VIS emission in the wavelength range from 400 to 700 nm.<sup>16</sup> Moreover, our recent study<sup>17</sup> showed that the  $Cr^{3+}/Yb^{3+}$  co-doped in the glasses generates emission spectra in the wavelength regions of 420–700 nm, 660–

c Faculty of Natural Sciences, Duy Tan University, Da Nang City, 550000, Vietnam

g Faculty of Applied Sciences, Ton Duc Thang University, Ho Chi Minh City, Vietnam

# Influences of copper–potassium ion exchange process on the optical bandgaps and spectroscopic properties of  $\text{Cr}^{3+}/\text{Yb}^{3+}$  co-doped in lanthanum

T. H. Le,<sup>a</sup> Anh-Luan Phan, <sup>bc</sup> Nguyen Minh Ty,<sup>d</sup> Dacheng Zhou,<sup>e</sup> Jianbei Qiu<sup>e</sup> a[n](http://orcid.org/0000-0002-0399-8422)d Ho Kim Dan **D**<sup>\*fg</sup>

aluminosilicate glasses

In this study, lanthanum aluminosilicate glasses with compositions of  $45SiO<sub>2</sub>-20Al<sub>2</sub>O<sub>3</sub>-12.5LaF<sub>3</sub>-10BaF<sub>2</sub>-$ 9K<sub>2</sub>O-1Cr<sub>2</sub>O<sub>3</sub>-2.5Yb<sub>2</sub>O<sub>3</sub> (SALBK) were prepared using the conventional melting method and copperpotassium ion exchange process. Influences of the ion exchange process between copper and potassium on the visible, upconversion, and near-infrared luminescence spectra of  $Cr<sup>3+</sup>/Yb<sup>3+</sup>$  co-doped under excitations of 343, 490, and 980 nm LD were investigated. The EDS analysis of SALBK glasses was measured to confirm the presence of atoms in the glasses. The values of direct and indirect bandgaps of  $Cr<sup>3+</sup>/Yb<sup>3+</sup>$  co-doped SALBK glasses were calculated and analyzed. Besides, the energy exchange processes between Cu<sup>+</sup>, Cu<sup>2+</sup> ions, and Cr<sup>3+</sup>, Yb<sup>3+</sup> ions were also proposed and discussed. **PAPER**<br>
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THE Lec<sup>2</sup> Anh-Luan Phan,<sup>16:</sup> Nguyen Minh Ty,<sup>2</sup> Dacheng Zhou,<sup>6</sup> Janbel Glu<sup>s</sup><br>
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860 nm, and 970–1150 nm corresponding to the excitations 358, 488, and 690 nm LD. Since then, we have been interested in enhancing emissions and optical properties of  $Cr^{3+}/Yb^{3+}$  codoped in the glasses. $17,18$  To this aim, embedding the coinage ions (such as,  $Ag^+$ ,  $Cu^+$  ions) into the glass through the ion exchange process $20-22$  is one of the different solutions which brought about positive results.<sup>3,12,19</sup> In fact, the ion exchange process between coinage ions and alkali ions has many advantages compared with other traditional methods.<sup>21</sup> For example, in many previous researches<sup>22-24</sup> as well as in our recent work,<sup>25</sup> it was shown that the coinage ions introduced by the ion exchange processes could significantly affect the optical properties of the glasses such as the refractive index, the optical density, the nearinfrared emission spectrum, or even chemically strengthen the glasses $26,27$  as well as modify glass structure.<sup>28</sup>

Through the ion exchange process, the Cu<sup>+</sup> and Cu<sup>2+</sup> ions, as well as  $Ag<sup>+</sup>$  ions, could be ejected into the surfaces of the glasses,22,24,29–<sup>32</sup> then become neutral copper or/and silver atoms and grow into copper or/and silver nanoparticles (CuNPs or/and AgNPs).<sup>24,29,32</sup> Therefore, the ion exchange process between copper or/and silver cations and alkali cations to enhance the luminescence of  $Cr^{3+}/RE^{3+}$  co-doped has been studied in recent times.<sup>21,29,33</sup> In this paper, we study the influences of the ion exchange process between copper and potassium on the optical bandgaps and spectroscopic properties of  $Cr^{3+}/Yb^{3+}$  co-doped in  $45SiO_2-20Al_2O_3-12.5LaF_3-10BaF_2-9K_2O-1Cr_2O_3-2.5Yb_2O_3$ lanthanum aluminosilicate glasses. We calculated the values of

both the direct and indirect optical bandgaps and figured out its manner of dependence on the salt concentration ratios between  $CuSO<sub>4</sub>:K<sub>2</sub>SO<sub>4</sub>$ . Besides, the energy transfer mechanism between

a Department of Physics and Technology, Thai Nguyen University of Sciences, Thai Nguyen, Vietnam

b Institute of Fundamental and Applied Sciences, Duy Tan University, Ho Chi Minh City 700000, Vietnam. E-mail: phananhluan@duytan.edu.vn

d Faculty of Natural Sciences, Thu Dau Mot University, Thu Dau Mot 590000, Vietnam e Key Laboratory of Advanced Materials of Yunnan Province, School of Materials Science and Engineering, Kunming University of Science and Technology, Kunming 650093, China

f Ceramics and Biomaterials Research Group, Advanced Institute of Materials Science, Ton Duc Thang University, Ho Chi Minh City, Vietnam. E-mail: hokimdan@tdtu.edu. vn

 $Cu<sup>+</sup>$  and  $Cu<sup>2+</sup>$  ions with  $Cr<sup>3+</sup>$ , Yb<sup>3+</sup> ions was also proposed and discussed.

### 2. Experimental details

In this study, we used the highly pure (99.99%) reagents of  $SiO<sub>2</sub>$ , Al<sub>2</sub>O<sub>3</sub>, LaF<sub>3</sub>, BaF<sub>2</sub>, Cr<sub>2</sub>O<sub>3</sub>, K<sub>2</sub>O, Yb<sub>2</sub>O<sub>3</sub>, K<sub>2</sub>SO<sub>4</sub>, and CuSO<sub>4</sub> to prepare compositions of  $45SiO<sub>2</sub>–20Al<sub>2</sub>O<sub>3</sub>–12.5LaF<sub>3</sub>–10BaF<sub>2</sub>–$  $9K_2O-1Cr_2O_3-2.5Yb_2O_3$  (SALBK) by the conventional melting method. After being condensed in a platinum crucible, 12 g of the mixture of these materials were put into an electric furnace to be heated at 1600 °C for 1 h under air atmosphere. After that, we poured the molten mixture into a mold placed on a polished plate made from stainless steel to form glass samples, then annealed all of them at 450  $^{\circ}$ C for 6 h to remove thermal strains.<sup>34</sup> We next cut them into 10 mm  $\times$  10 mm  $\times$  2 mm size and finally polished their surface for the sake of measurements.

In order to perform the ion exchange process between copper and potassium, we prepared salt mixtures with different concentration ratios between  $xCuSO_4$ :(100 - x)K<sub>2</sub>SO<sub>4</sub> where x varies as particularly given in Table 1.

All the experimental measurements with the glass samples, which were carried out at the ambient air temperature, and their corresponding instruments are listed in Table 2.

To find the suitable temperature ranges for the ion exchange process between copper and potassium, we conducted DTA analysis for  $45SiO_2-20Al_2O_3-12.5LaF_3-10BaF_2-9K_2O-1Cr_2O_3 2.5Yb<sub>2</sub>O<sub>3</sub>$  (SALBK) glass sample. The results are shown in Fig. 1, where  $T_{\rm g}$ ,  $T_{\rm x}$ , and  $T_{\rm p}$  are the glass transition temperature, the crystallization onset temperature and crystallization peak temperature, respectively. According to the results, we chose 587 °C as the temperature of the salt mixtures into which we submerged the glass samples for 24 h.<sup>34</sup>



Fig. 1 DTA analysis of  $45SiO<sub>2</sub> - 20Al<sub>2</sub>O<sub>3</sub> - 12.5LaF<sub>3</sub> - 10BaF<sub>2</sub> - 9K<sub>2</sub>O 1Cr_2O_3-2.5Yb_2O_3$  (SALBK) glass sample.

Next, they were taken out and washed with alcohol and distilled water to eliminate all the residual salt on their surfaces. After all, the glass samples were further heat-treated at 686  $\degree$ C for 8 h to promote the copper nanoparticles (CuNPs) formation.

# 3. Results and discussion

The EDS analysis of the SALBK-1Cr2.5Yb-35Cu glass sample is shown in Fig. 2. In addition to the EDS peaks of the Si, O, Al, La, F, Ba, K, Cr, and Yb atoms in SALBK glass,<sup>35</sup> the ones of copper were strongly obtained corresponding to the energy values about 0.95, 8.04, and 8.99 keV, $36$  which means that the copper and potassium ions have been added into the glass network through the ion exchange process.<sup>37</sup>

Table 1 The concentration ratio of salts mixture for the ion exchange process



Table 2 The measurements and their corresponding instruments





Fig. 2 EDS analysis of SALBK-1Cr2.5Yb-35Cu glass sample.



Fig. 3 XRD analysis of SALBK-1Cr2.5Yb-0Cu, SALBK-1Cr2.5Yb-15Cu, SALBK-1Cr2.5Yb-20Cu, SALBK-1Cr2.5Yb-25Cu, SALBK-1Cr2.5Yb-30Cu, and SALBK-1Cr2.5Yb-35Cu glass samples.



Fig. 4 Absorption spectra of SALBK-1Cr2.5Yb-0Cu, SALBK-1Cr2.5Yb-15Cu, SALBK-1Cr2.5Yb-20Cu, SALBK-1Cr2.5Yb-25Cu, SALBK-1Cr2.5Yb-30Cu, and SALBK-1Cr2.5Yb-35Cu glass samples.



Fig. 5 XPS spectra of SABLK-15Cu glass sample.

To investigate the influence of the ion exchange process between copper and potassium on the structure of glass materials, we performed the XRD analysis of all the glass samples together with using X'Pert HighScore Plus software,<sup>38</sup> showing the results in Fig. 3. Conspicuously, the XRD pattern of the SALBK-1Cr2.5Yb-0Cu glass sample has no diffraction peak,<sup>39</sup> while all of the others show three main peaks at  $2\theta = 30.6$ degree (110),  $2\theta = 44.2$  degree (111) and  $2\theta = 52.1$  degree (200)<sup>40</sup> due to the formation of copper nanoparticles (CuNPs).<sup>41</sup> Besides, no diffraction peak of other nanocrystals is observed for all the samples.

Next, we exhibit the absorption spectra of SALBK-1Cr2.5Yb-0Cu, SALBK-1Cr2.5Yb-15Cu, SALBK-1Cr2.5Yb-20Cu, SALBK-1Cr2.5Yb-25Cu, SALBK-1Cr2.5Yb-30Cu, and SALBK-1Cr2.5Yb-35Cu glass samples in Fig. 4. For the SALBK-1Cr2.5Yb-0Cu glass sample, we can observe three absorption peaks bands centered at 470, 650, and 878 nm, which is evidence that they are from the  ${}^4\text{A}_{2g} \rightarrow {}^4\text{T}_{1g}(\text{F})$ ,  ${}^4\text{A}_{2g} \rightarrow {}^4\text{T}_{2g}(\text{F})$  and  ${}^4\text{A}_{2g} \rightarrow {}^2\text{E}$ transitions of  $Cr^{3+}$ ,  $17,42,43$  In addition, with the increase in salt concentration ratio  $p$  from 0.18 to 0.54, the emission intensities of  $Cr^{3+}/Yb^{3+}$  co-doped bands centered at 470, 650 nm, and



Fig. 6 Visible emission spectra of SALBK-1Cr2.5Yb-0Cu, SALBK-1Cr2.5Yb-15Cu, SALBK-1Cr2.5Yb-20Cu, SALBK-1Cr2.5Yb-25Cu, SALBK-1Cr2.5Yb-30Cu, and SALBK-1Cr2.5Yb-35Cu glass samples under excitation 343 nm.

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Name of glass samples	CIEx	CIEV	Position on the CIE $1931(x, y)$ chromaticity coordinates	Color region
SALBK-1Cr2.5Yb-0Cu	0.2511	0.4241	P <sub>0</sub> Cu	Green
SALBK-1Cr2.5Yb-15Cu	0.2648	0.4356	P <sub>15</sub> Cu	Green
SALBK-1Cr2.5Yb-20Cu	0.2562	0.4294	P <sub>20</sub> Cu	Green
SALBK-1Cr2.5Yb-25Cu	0.2722	0.4363	P <sub>25</sub> Cu	Green
SALBK-1Cr2.5Yb-30Cu	0.2779	0.4402	P30Cu	Green
SALBK-1Cr2.5Yb-35Cu	0.2851	0.4501	P35Cu	Yellowish-green
to the local surface plasmon resonance (LSPR) of copper ions <sup>44</sup> and the crystal field variation caused by copper ions. <sup>17,43,45</sup> Moreover, the intensities of two absorption spectra bands centered at 306 nm and 878 nm also increased significantly, which confirms the existence and the role of $Cu^{+}$ and $Cu^{2+}$ , respectively, <sup>46</sup> in the absorption spectra of $Cr^{3+}/Yb^{3+}$ co-doped.		878 nm strongly increased. These increments may be ascribed	both Cu <sup>+</sup> and Cu <sup>2+</sup> ions in the SALBK glasses, corresponding to the XPS major peaks at 932.39 eV and 934.91 eV. <sup>49-52</sup> In Fig. 6, we present the visible emission spectra under excitation 343 nm of all these glass samples with two peaks at about 488 and 653 nm, which are assigned to the ${}^{2}T_{2} \rightarrow {}^{4}A_{2g}$ and ${}^{4}T_{2g}({}^{4}F) \rightarrow {}^{4}A_{2g}$ transitions of Cr <sup>3+</sup> . <sup>17,43,45</sup> As shown, the visible emission intensities of $Cr^{3+}/Yb^{3+}$ co-doped bands	
The Cu <sup>+</sup> formed and exists in silicate glass due to reducing of $Cu^{2+47,48}$ To further demonstrate the existence of both $Cu^{+}$ and $Cu2+$ ions in the glass after ion exchange. We analyzed XPS spectra of the SABLK-15Cu glass sample, the results are shown in Fig. 5 (the blue and green curves are the curve-fitting Gaussian of XPS spectra at 934.91 eV and 932.39 eV, respec- tively). Based on these results, we can confirm the existence of			centered at 488 and 653 nm significantly increased with the increase in salt concentration ratio $p$ from 0.18 to 0.54. <sup>47</sup> Interestingly, the peak at 488 nm of $Cr^{3+}$ under 343 nm excita- tion has a slight blue-shift of about 8 nm. The emission peak shift at 488 nm of $Cr^{3+}$ is assigned to the role of $Cu^{+}$ cations. This result is also consistent with the suggestions discussed in the paper <sup>53</sup> of Tian-Shuai Lv et al.	

Table 3 CIE 1931 (x; y) chromaticity coordinates for luminescence of  $Cr^{3+/Yb^{3+}}$  co-doped in SALBK-1Cr2.5Yb-0Cu, SALBK-1Cr2.5Yb-15Cu, SALBK-1Cr2.5Yb-20Cu, SALBK-1Cr2.5Yb-25Cu, SALBK-1Cr2.5Yb-30Cu, and SALBK-1Cr2.5Yb-35Cu glass samples



Fig. 7 CIE 1931(x; y) chromaticity coordinates for luminescence of  $Cr^{3+}/Yb^{3+}$  co-doped in SALBK-1Cr2.5Yb-0Cu, SALBK-1Cr2.5Yb-15Cu, SALBK-1Cr2.5Yb-20Cu, SALBK-1Cr2.5Yb-25Cu, SALBK-1Cr2.5Yb-30Cu, and SALBK-1Cr2.5Yb-35Cu glass samples.



Fig. 8 UC spectra of  $Cr^{3+}/Yb^{3+}$  co-doped in SALBK-1Cr2.5Yb-0Cu, SALBK-1Cr2.5Yb-15Cu, SALBK-1Cr2.5Yb-20Cu, SALBK-1Cr2.5Yb-25Cu, SALBK-1Cr2.5Yb-30Cu, and SALBK-1Cr2.5Yb-35Cu glass samples under excitation 980 nm LD.

The increases in visible emission intensity of  $Cr^{3+}/Yb^{3+}$  codoped band centered at 488 nm and 653 nm were assigned to the energy transfer (ET<sub>1</sub>) process from  $^3\text{E}_\text{g}\text{(T_{2g})} \rightarrow \, ^1\text{A}_\text{g}$  transition of Cu<sup>+</sup> to <sup>2</sup>T<sub>2</sub>  $\rightarrow$  <sup>4</sup>A<sub>2g</sub> transition of Cr<sup>3+</sup>,<sup>47,48</sup> and to the ET<sub>2</sub> process from  $^3\text{E}_\text{g}(\text{T}_{1 \text{g}}) \rightarrow \,^1\text{A}_\text{g}$  transition of Cu $^+$  to  $^4\text{T}_{2 \text{g}}(^4\text{F}) \rightarrow \,^4\text{A}_{2 \text{g}}$ transition of Cr $^{3+}, ^{47,48}$  respectively. These two ET $_1,$  ET $_2$  processes are proposed as follows:

 $ET_1: {}^3E_g(T_{2g})(Cu^+) + {}^4A_{2g}(Cr^{3+}) \rightarrow {}^2T_2(Cr^{3+}) + {}^1A_g(Cu^+).$  $ET_2$ :  ${}^{3}E_g(T_{1g})(Cu^+) + {}^{4}A_{2g}(Cr^{3+}) \rightarrow {}^{4}T_{2g}({}^{4}F)(Cr^{3+}) + {}^{1}A_g(Cu^+).$ 

Furthermore, the calculated results of the CIE 1931 $(x; y)$ chromaticity coordinates for luminescence of  $Cr^{3+}/Yb^{3+}$  codoped in SALBK-1Cr2.5Yb-0Cu, SALBK-1Cr2.5Yb-15Cu, SALBK-1Cr2.5Yb-20Cu, SALBK-1Cr2.5Yb-25Cu, SALBK-1Cr2.5Yb-30Cu, and SALBK-1Cr2.5Yb-35Cu glass samples in correspondence to the P0Cu, P15Cu, P20Cu, P25Cu, P30Cu, and P35Cu points on the CIE 1931 $(x, y)$  chromaticity coordinates are given in Table 3, while Fig. 7 is for the sake of illustration. It can be seen that except that the CIE 1931 $(x; y)$  chromaticity coordinates for luminescence of Cr3+/Yb3+ co-doped in SALBK-1Cr2.5Yb-35Cu



Fig. 9 NIR emission spectra of  $Cr^{3+}/Yb^{3+}$  co-doped in SALBK-1Cr2.5Yb-0Cu, SALBK-1Cr2.5Yb-15Cu, SALBK-1Cr2.5Yb-20Cu, SALBK-1Cr2.5Yb-25Cu, SALBK-1Cr2.5Yb-30Cu, and SALBK-1Cr2.5Yb-35Cu glass samples, excited by 490 nm.

glass sample were shifted to the yellowish-green color region, the remaining were originally in the green color region.

Fig. 8 shows the upconversion (UC) emission spectra of  $Cr^{3+}/$  $Yb^{3+}$  co-doped in all the glass samples under excitation 980 nm LD with two bands centered at  $\sim$ 537 and 853 nm corresponding to  ${}^{2}T_{2}$   $\rightarrow$   ${}^{4}A_{2g}$ , and  ${}^{2}E$   $\rightarrow$   ${}^{4}A_{2g}$  transitions of Cr<sup>3+</sup>, respectively.<sup>17,45,54-56</sup> With the increase in salt concentration ratio  $p$ from 0.18 to 0.54, the emission intensities of these bands markedly increased. These increments were assigned to the aforementioned  $ET_1$  process (for the band centered at 537 nm) and to the ET<sub>3</sub> process from  ${}^{2}B_{2g} \rightarrow {}^{2}B_{1g}$  transition of Cu<sup>2+</sup> to  ${}^{2}E$  $\rightarrow$  <sup>4</sup>A<sub>2g</sub> transition of Cr<sup>3+</sup> (for the one at 853 nm),<sup>5,46</sup> which is proposed as follows:

 $ET_3$ :  ${}^2E_g(Cu^{2+}) + {}^4A_{2g}(Cr^{3+}) \rightarrow {}^2E(Cr^{3+}) + {}^2B_{1g}(Cu^{2+}).$ 

Likewise, Fig. 9 shows the NIR emission spectra under excitation 490 nm. There was only one NIR emission peak at 1016 nm, and it is assigned to the <sup>2</sup> $F_{5/2}$   $\rightarrow$  <sup>2</sup> $F_{7/2}$  transition of Yb<sup>3+</sup>.<sup>3,5</sup> Similarly, the NIR emission intensity of the band centered at 1016 nm also increased with the increase in salt concentration ratio p, which is a manifestation of the  $ET<sub>4</sub>$ process from <sup>2</sup> $B_{2g} \rightarrow {}^{2}B_{1g}$  transition of Cu<sup>2+</sup> to <sup>2</sup> $F_{5/2} \rightarrow {}^{2}F_{7/2}$ transition of Yb<sup>3+</sup>:<sup>57</sup>

 $ET_4$ :  ${}^{2}E_g(Cu^{2+}) + {}^{2}F_{7/2}(Yb^{3+}) \rightarrow {}^{2}F_{5/2}(Yb^{3+}) + {}^{2}B_{1g}(Cu^{2+}).$ 

The mechanism for the visible and NIR luminescence of  $Cr^{3+}/Yb^{3+}$  co-doped under excitations of 343, 490, and 980 nm LD, and the details of all the above-mentioned  $ET_1$ ,  $ET_2$ ,  $ET_3$ , and  $ET_4$  processes are described in Fig. 10.



Fig. 10 Mechanism ET processes for the visible, UC, and NIR luminescence of  $Cr^{3+}/Yb^{3+}$  co-doped under excitations 343, 490, and 980 nm LD.



Fig. 11 (A) Decay lifetimes curves of Cr<sup>3+</sup> at 537 nm in SALBK-0Cu, SALBK-15Cu, SALBK-20Cu, SALBK-25Cu, SALBK-30Cu, and SALBK-35Cu glass samples, under excitation 980 nm. (B) Decay lifetimes curves of Cr<sup>3+</sup> at 653 nm in SALBK-0Cu, SALBK-15Cu, SALBK-20Cu, SALBK-25Cu, SALBK-30Cu, and SALBK-35Cu glass samples, under excitation 343 nm. (C) Decay lifetimes curves of Cr3+ at 853 nm in SALBK-0Cu, SALBK-15Cu, SALBK-20Cu, SALBK-25Cu, SALBK-30Cu, and SALBK-35Cu glass samples, under excitation 980 nm LD. (D) Decay lifetimes curves of Yb<sup>3+</sup> at 1016 nm in SALBK-0Cu, SALBK-15Cu, SALBK-20Cu, SALBK-25Cu, SALBK-30Cu, and SALBK-35Cu glass samples, under excitation 490 nm.

#### Table 4 Decay lifetimes values of  $Cr^{3+}$  and Yb<sup>3+</sup> demonstrating for the ET<sub>1</sub>, ET<sub>3</sub>, ET<sub>3</sub>, and ET<sub>4</sub> processes



To further validate evidence for the  $ET_1$ ,  $ET_2$ ,  $ET_3$ ,  $ET_4$ processes from Cu $^{\dagger}/\text{Cu}^{2+}$  ions to Cr $^{3+}$  and Yb $^{3+}$  ions, we carried out the decay lifetimes measurement for several glass samples. Namely, the decay lifetimes curves of  $Cr^{3+}$  at 537, 653, and 853 nm to demonstrations for the  $ET_1$ ,  $ET_2$ ,  $ET_3$  processes are shown in Fig. 11A–C, respectively while the decay lifetimes curve of Yb<sup>3+</sup> at 1016 nm to demonstrations for the  $ET_4$  is in Fig. 11D.

We recall that the decay lifetimes can be calculated by the following equation:58,59

$$
\tau = \frac{A_1 \tau_1^2 + A_2 \tau_1^2}{A_1 \tau_1 + A_2 \tau_1}
$$
 (1)

where  $A_1, A_2$  are constants;  $\tau_1, \tau_2$  are rapid and slow lifetimes for exponential components, respectively. Using the formula (1) for the data obtained in Fig. 11A–D, we calculated the decay lifetime values of  $Cr^{3+}$  and  $Yb^{3+}$ , corresponding to the  $ET_1, ET_2, ET_3$ and  $ET_4$  processes, details are presented in Table 4.

Finally, we calculated and analyzed the optical bandgap  $(E_{\varphi})$ of  $Cr^{3+}/Yb^{3+}$  co-doped in SALBK glasses to evaluate whether and how it is affected by the ion exchange process between copper and potassium. The optical bandgap  $E_g$  can be calculated using Tauc's formula:<sup>60</sup>

$$
\alpha h\nu = A(h\nu - E_{g})^{m} \tag{2}
$$

where  $\alpha$  is the absorption coefficient, A is a constant, m is the power depending on the nature of transition ( $m = 1/2$  for direct transition, and  $m = 2$  for indirect transition).<sup>61-63</sup> From formula (2), the values of both types (direct and indirect) of the optical bandgap in SALBK glasses are calculated and detailed in Table 5. For the sake of demonstration, we also present the plot of  $(hv)$ *versus* both  $(\alpha h\nu)^2$  and  $(\alpha h\nu)^{1/2}$  for estimating  $E_{\rm g}$  of all the considered samples in Fig. 12 and 13.

The calculation gave us the estimated results of 3.23–3.33 eV for direct bandgap and 2.83–3.02 eV for indirect bandgap. On the other hand, both bandgap types decrease when the salt concentration ratio  $p$  increases from 0.18 to 0.54. Thus, we can conclude that the ion exchange process between copper and potassium had a diminishing effect on the bandgap. The reasons for this effect can be: (i) Because there are non-bridging oxygens (NBOs) in the silicate glasses network,<sup>64</sup> the  $Al^{3+}$  ions have some options to link with SiO<sub>4</sub> tetrahedra to form (Al, Si)-O-Si, (Al, Si)-O-Al bonds,<sup>65</sup> or with others groups to form (Al, Si)–O bonds.<sup>65</sup> After the copper ions were introduced into the

Table 5 The values of both the direct and indirect optical bandgaps in SALBK glasses

Glass samples	Direct bandgap $hv$ [(eV)]	Indirect bandgap $hv$ [(eV)]
SALBK-1Cr2.5Yb-0Cu	3.33	3.02
SALBK-1Cr2.5Yb-15Cu	3.32	2.98
SALBK-1Cr2.5Yb-20Cu	3.30	2.91
SALBK-1Cr2.5Yb-25Cu	3.28	2.88
SALBK-1Cr2.5Yb-30Cu	3.25	2.87
SALBK-1Cr2.5Yb-35Cu	3.23	2.83



Fig. 12 Plot of  $(hv)$  versus  $(\alpha h v)^2$  for estimating the  $E_g$  of SALBK-1Cr2.5Yb-0Cu, SALBK-1Cr2.5Yb-15Cu, SALBK-1Cr2.5Yb-20Cu, SALBK-1Cr2.5Yb-25Cu, SALBK-1Cr2.5Yb-30Cu, and SALBK-1Cr2.5Yb-35Cu glass samples.



Fig. 13 Plot of  $(hv)$  versus  $(\alpha h v)^{1/2}$  for estimating the  $E_g$  of SALBK-1Cr2.5Yb-0Cu, SALBK-1Cr2.5Yb-15Cu, SALBK-1Cr2.5Yb-20Cu, SALBK-1Cr2.5Yb-25Cu, SALBK-1Cr2.5Yb-30Cu, and SALBK-1Cr2.5Yb-35Cu glass samples.

silica network by the copper-potassium ion exchange process, these bonds may be broken and then  $Al-O^-$  and  $Si-O^-$  groups can combine with copper ions to create the Si–O–Cu, and Al–O– Cu bonds. (ii) With the increase of salt concentration ratio, more negative sites appeared due to the local structure of the NBOs in the silicate glasses network.<sup>64,66</sup>

### 4. Conclusions

The CuNPs were formed in  $45SiO_2-20Al_2O_3-12.5LaF_3-10BaF_2 9K_2O-1Cr_2O_3-2.5Yb_2O_3$  lanthanum aluminosilicate glasses through the ion exchange process between copper and potassium process. The intensities of all three visible, UC, and NIR emissions of  $Cr^{3+}/Yb^{3+}$  co-doped bands at 480, 537, 653, 853, and 1016 nm increased with the increase in the ratio of salt concentrations  $CuSO<sub>4</sub>:K<sub>2</sub>SO<sub>4</sub>$  from 0.18 to 0.54. When the salt concentration ratio of the ion exchange process achieved 35 mol%CuSO<sub>4</sub>:65 mol%K<sub>2</sub>SO<sub>4</sub>, the CIE 1931(x; y) chromaticity

coordinates for the luminescence of  $Cr^{3+}/Yb^{3+}$  co-doped in SALBK-1Cr2.5Yb-35Cu glass sample shifted to the yellowishgreen color region. At the same time, the estimated results of the optical bandgap  $(E_g)$  confirmed that with the increase in the ratio of salt concentrations  $CuSO<sub>4</sub>:K<sub>2</sub>SO<sub>4</sub>$  from 0.18 to 0.54, the value of both direct and indirect bandgaps decrease. Besides, the possible energy transfer processes from  $Cu^+$ ,  $Cu^{2+}$  ions to  $Cr^{3+}$ , Yb<sup>3+</sup> ions were determined through the experimental results of the spectroscopic emissions and the decay lifetimes. RSC Advances Continuus contents for the Licence of C<sup>2+</sup>/15<sup>2</sup> codepted on 26 February 2021. Downloaded on 26 February 2021. Due to the first content of the spectroscopic material commons are the spectroscopic interest an

# Conflicts of interest

There are no conflicts to declare.

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