A new overview on rare earth diphenylphosphinates: Europium characteristic luminescence of low dimensional nanostructured materials

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1. Introduction

The search for highly porous and crystalline polymer compounds has become relevant in the research around the world and has generated, in the last three decades [1], a growing interest not only in the scientific community but also in several areas of technology. Main reasons for this are due to their supramolecular structures, aesthetically attractive structural topologies and potentialities for innovative technological applications. However, wet route precipitation or hydro/solvothermal methods may lead to the formation of highly insoluble porous coordination networks, as is true of europium diphenylphosphinates – \([\text{Eu}(\text{dpp})]_n\). The aim of this work was to study the optical behavior and its relation to the structure of \([\text{Eu}(\text{dpp})]_n\) complexes obtained through different synthetic approaches by precipitation: wet route (WRS), ultrasound-assisted wet route (U-WRS), modulator-assisted wet route (M-WRS) and wet route synthesis with solvothermal treatment (S-WRS). These methodologies allowed to synthetize crystalline materials in which both chelate and bridged coordination modes coexist, and they were efficient for obtaining stick-shaped 1D1 nanostructured materials with well-defined optical properties. Changes in precipitation kinetics and crystalline phases formation rate led to a distribution of \(\text{Eu}^{3+}\) ions in higher symmetry sites (HSS). S-WRS, transforming part of the less symmetrical sites into HSS, provided a better ordering of \([\text{Eu}(\text{dpp})]_n\). Furthermore, this is the first report of two or more symmetry sites for \(\text{Eu}^{3+}\) in a pure \([\text{Eu}(\text{dpp})]_n\) matrix.

These trivalent rare earth-based MOFs – RE³⁺MOFs – [11] present odd luminescent and magnetic properties. This is due to the different RE³⁺-RE³⁺ and/or RE³⁺-ligand chemical environments imposed by the coordination network, leading to several relaxation mechanisms, such as ligand-RE³⁺ cross relaxation and 4f-4f intraconfigurational transitions.

Among the several reported applications in the literature for RE³⁺ MOFs, it is possible to highlight their use in the sensing of humidity, pH, temperature and explosives [12,13]. Therefore, it is of interest the rational design and synthesis of RE³⁺ MOFs, looking forward to a better understanding of their optical and structural properties, which could allow applications in an improved way with great technological power.

However, obtaining RE³⁺ MOFs by wet route precipitation or
hydro/solvothermal methods may lead to the formation of highly insoluble porous coordination networks in different solvents commonly used in the laboratory, either polar or non-polar.

This variant makes it difficult to obtain crystals with a long-range periodic arrangement of perfectly stacked unit cells that can be easily characterized by single crystal X-ray diffraction (SCXRD), considered one of the most powerful techniques for structural elucidation of crystalline materials [14].

Rare earth diphenylphosphinates – [RE(dpp)]n, synthesized by direct reaction between a rare earth chloride (RECl3) and diphenylphosphinic acid (Hdpp), are authentic examples that integrate this class of materials presenting very low solubilities that do not produce single crystals.

In this work, we report for the first time – to the best of our knowledge – that modifications in synthetic methods and strict control of synthesis parameters in [RE(dpp)]n preparations could lead to the formation of materials with a greater crystallinity with defined morphology and size and mainly with well-defined optical properties, opening opportunities for future technological applications of these phosphinates in lighting, ionizing radiation detection and sensing.

To achieve that, four different synthetic approaches by precipitation were addressed: wet route (WRS), ultrasound-assisted wet route (U-WRS), modulator-assisted wet route (M-WRS) and wet route synthesis with solvothermal treatment (S-WRS).

Acoustic cavitation in [RE(dpp)]n synthesis aims the design of an environment by which the intercurrence of homogeneous nucleation and a significant increase in the speed of recrystallization are prioritized when compared to the conventional methods applied in the preparation of nanostructured materials [15–18].

On the other hand, coordination modulators behave in a competitive way with the organic ligand by coordination with metallic ions present in solution, regulating nucleation and reducing crystal growth rate and avoiding precipitation of amorphous products [10,19]. In this way, they provide a more accurate control of the size and morphology of the crystals, allowing the formation of nanoparticles [20]. Indeed, modulation is a suitable strategy to improve the final product crystallinity and the synthesis reproducibility [21].

And last but not least, solvothermal process is useful for achieving thermodynamically stable and metastable states, contributing to the formation of new materials which cannot be easily synthesized from conventional synthetic routes [22].

1.1. Background

Studied since the 1970s, [RE(dpp)]n, are presented as peculiar and somewhat intriguing materials. They have two distinct symmetry sites for trivalent rare earth, thermal stability similar to that of inorganic phosphates and mesoporous structure with the presence of crystalline and non-crystalline domains.

In the first report of rare earth diphenylphosphinates preparation, Dunstan and Vicentini [23] proposed the general formula RE(dpp)3.n (H2O) for the synthesized coordination compounds.

Bel’tyukova [24] in 1989 published the first spectroscopic study involving compounds formed by trivalent europium and diphenylphosphinate ligand reported in the literature.

Innocentini and co-workers [25] synthesized a non-hygroscopic europium diphenylphosphinate with high thermal stability in comparison with other coordination compounds and reasonable ligand-to-metal energy transfer ability via antenna effect.

Stucchi [26], Scarpari [27,28] and their collaborators prepared a series of rare earth diphenylphosphinates and binary compounds, examining their optical behavior to predict some structural information.

Francisco et al. [29] suggested the presence of one or more symmetry sites for RE3+ in a europium-doped yttrium diphenylphosphinate matrix. In another study [30], applying the hydrothermal synthesis, they obtained unidimensional structures in the form of rods.

According to Abreu et al. [31,32], [RE(dpp)]n are polycrystalline materials in a polymeric arrangement, resulting in coordination sites of different symmetries, and can be represented by [RE(dpp)]n.

Souza [33], Munhoz [34] and their co-workers have investigated the potential application of lanthanum and cerium diphenylphosphinates (and their doping) as UV absorbers.

Rosa et al. [35] examined the effect of cerium as a sensitizer and activator on the compounds [La(dpp)]3:Eu3+ and [La(dpp)]3:Eu3+,Tb3+ and their possible uses as emission-emitting luminescent materials of white light.

In addition, Rosa et al. [36] proposed that diphenylphosphinate coordination compounds of La3+-Ce3+-Eu3+ / La3+-Ce3+-.Tb3+ ternary systems exist as cross-linked coordination polymers, forming a bimolecular system: [Eu3(dpp)]n. They also verified that those ternary systems have an appreciable resistance to ionizing radiation.

Recently, Bim et al. [37] carried out a systematic study of trivalent rare earth diphenylphosphinates in an attempt to elucidate the structure and symmetry of the trivalent lanthanide crystallographic site, as well as the coordination mode of the dpp ligand to RE3+ cations.

Many gaps regarding the structure of [RE(dpp)]n are still found without complete investigation data. In previously presented works, the authors did not include details about the composition and coordination sphere of the compounds, nor did they refer to the symmetry of RE3+ crystallographic site, the coordination mode of dpp ligand to metal cations or thermal behavior presented by them.

Thus, a more comprehensive study of rare earth diphenylphosphinates in terms of their structures and intrinsic optical properties will provide the means to both promote them to the level of metal-organic frameworks and to suggest applications of scientific, technological and industrial interest.

2. Experimental section

[RE(dpp)]n synthesis was reviewed and modified from that initially proposed by Stucchi et al. [26] to more precise and controlled synthesis parameters, such as temperature and atmosphere of the reaction medium, stirring, reaction time and solvent used. It is emphasized that the authors of this paper have been working with the complete series of rare earth diphenylphosphinates ([RE(dpp)]n, RE3+ = Y3+, La3+, Ce3+, Eu3+, Tb3+, Nd3+) or their doping; however, for the scope of this paper, we decided to focus on [Eu(dpp)]n, results, addressing the intrinsic optical properties of trivalent europium ion. Purity and brands: EuCl3 solution was obtained by the stoichiometric hydrochloric acid (Sigma-Aldrich, 37%) attack on Eu2O3 (Aldrich, 99.99%) and its concentration was determined by complexation titrimetry, diphenylphosphinic acid (Sigma-Aldrich, ≥ 98%) attack on Eu2O3 (Alrich, 99.99%) and its concentration was determined by complexation titrimetry, diphenylphosphinic acid (Alrich, ≥ 98.0%), methanoic acid (Sigma-Aldrich, ≥ 95%), ethanol (Sigma-Aldrich, ≥ 99.5%), compressed nitrogen gas (White Martins, 99.998%).

2.1. Wet route precipitation synthesis (WRS)

The preparation consisted of mixing EuCl3 solution (4.40 mL, 0.4488 mmol) with ligand solution (diphenylphosphinic acid, Hdpp – 0.2998 g, 1.3464 mmol) in 1:3 metal-to-ligand ratio. The solvent used was 40.00 mL of a 1:1 mixture of ethanol and distilled water. The ligand solution was placed in a three-necked round bottom flask, kept in a microprocessed hot water bath. The temperature of the reaction medium was maintained at 72 ± 3°C and measured by thermocouple (NTC sensor - nickel-plated brass - Ø6.35 mm/L = 22 mm) coupled to a digital temperature controller (Ageon G101 Color). The ligand solution was subjected to constant stirring and the round bottom flask saturated with N2(g). EuCl3 solution was slowly dripped onto the ligand solution via a 10 mL burette, causing the precipitation of [Eu(dpp)]n. After the addition of EuCl3 solution, the mixture (precipitate and supernatant) remained in the hot water bath under stirring for 60 min. The flask was then removed from the system and kept under stirring for extra 30 min.

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At the end of this process, the contents of the flask (precipitate and supernatant) were transferred to a 50-mL centrifuge tube, remaining at rest for 24 h. After this period, the mixture was centrifuged at 10,000 rpm for 10 min (Eppendorf Centrifuge 5804R). The precipitate was separated from the supernatant and washed with the ice-cold ethanol/water (1:1) mixture, then settled in a desiccator containing silica gel for 7 days.

2.2. Ultrasound-assisted wet route precipitation synthesis (U-WRS)

The synthesis of ultrasound-assisted europium diphenylphosphinate was carried out in a cell disruptor (R2D091109 - UNIQUÉ). It consisted of mixing EuCl₃ solution (4.40 mL, 0.4488 mmol) with Hdpp solution (0.2998 g, 1.346 mmol) in 1:3 metal-to-ligand ratio subjected to high intensity sonication. The solid ligand was placed in a three-necked round bottom flask containing 40.00 mL of solvent (a 1:1 mixture of ethanol and distilled water). This flask was kept immersed in a bath with water at room temperature and continuously saturated with N₂(g). A titanium horn driven by a piezoelectric transducer was introduced directly into the solution through the central mouth of the round bottom flask and 20 kHz pulses were applied for 2 min leading to complete solubilization of diphenylphosphinic acid. Keeping the pulse sequence, EuCl₃ solution was slowly dripped onto Hdpp solution by means of a 10-mL burette, causing the precipitation of [Eu(dpp)]₃n. After the addition of EuCl₃ solution, the mixture (precipitate and supernatant) remained under cavitation for 8 min. At the end of this process, the contents of the flask (precipitate and supernatant) were transferred to a 50-mL centrifuge tube, remaining at rest for 24 h. The other procedures were identical to those described at the end of Subsection 2.1.

2.3. Modulator-assisted wet route precipitation synthesis (M-WRS)

For the synthesis of modulator-assisted europium diphenylphosphinate, we used a monocarboxylic acid as modulating agent, more specifically, methanoic acid. The whole procedure is similar to those described in Subsection 2.1, only adding the methanoic acid (1.82 mL, 44.88 mmol) to the flask containing the already solubilized ligand.

2.4. Wet route precipitation synthesis with solvothermal treatment (S-WRS)

The preparation of [Eu(dpp)]₃n was carried out as described in Subsection 2.1 until prior to its centrifugation. The contents of the flask (precipitate and supernatant) were transferred to a Teflon beaker, keeping 30% of its useful volume unoccupied. N₂(g) was used to saturate the medium. The Teflon beaker was then introduced into a stainless-steel reactor, arranged in a circulation oven (Model MSM 510/27/CR/ RP, Series 1105.0218 – M.S. Mistura Equipamentos para Laboratório) and subjected to controlled temperature programming: heating rate, 3.2 °C h⁻¹; landing, 180 °C; 48 h; cooling rate up to room temperature, 1.6 °C h⁻¹. After the solvothermal treatment, the reactor contents (precipitate and supernatant) were transferred to a 50-mL centrifuge tube, remaining at rest for 24 h. The other procedures were identical to those described at the end of Subsection 2.1.

2.5. Characterisation

[Eu(dpp)]₃n obtained through the four routes described above (Subsections 2.1–2.4) have been subjected to several structural and spectroscopic characterization techniques. Elemental analysis was performed on a Perkin Elmer 2400 Series II CHNS/O Elemental Analyzer. Analysis based on X-ray diffraction was carried out on Siemens diffractometer with copper rotational anode using Cu-Kα1 radiation (1.5418 Å) and graphite curved monochromator. [Eu(dpp)]₃n morphology and size were observed by using scanning electron microscope (SEM) JEOL JSM7500F equipped with a field emission gun (FEG) operated at 2.00 kV; particle counting and determination of their dimensions were performed by inspection with the aid of an image processing and analysis software in Java, ImageJ. Photoluminescence emission and excitation spectra were obtained using a Horiba Jobin Yvon spectrophotometer, model Fluorolog-3, and registered in UV–Vis region using a photomultiplier Horiba Jobin Yvon R928 and xenon lamp (ozone free); emission spectra were collected in triplicate and submitted to one-way ANOVA (analysis of variance) with post-hoc Tukey HSD (honestly significant difference) test. Vibrational spectra were obtained with BRUKER Fourier-transform infrared spectrophotometer - VERTEX 70 with DLaTGS detector, using an attenuated total reflectance (ATR) accessory.

3. Results and discussion

The new methodologies (Subsections 2.1–2.4) adopted for [Eu(dpp)]₃n preparation had positive impact on the precipitation of the metal complexes, providing controlled growth of materials remarkably more crystalline and organized than those previously reported in the literature [26,27,30,36] (Fig. 1).

Independently of the adopted synthetic route, the precipitations were stoichiometric; the overall reactions yields were always higher than 90% and the losses are related to the processes of washing, transference and maceration of precipitates. Elemental analyzes of [Eu(dpp)]₃n synthesized by WRS, U-WRS, M-WRS and S-WRS provided carbon, hydrogen and nitrogen (adsorbed in the pores of the structures) contents ranging between 54.84–55.10%, 3.66–3.78% and 2.12–2.82%, respectively.

XRD pattern of WRS [Eu(dpp)]₃n is shown in Fig. 1(a) and should be regarded as representative of whole synthesized series through the synthetic modifications proposed by the authors of this paper; U-WRS, M-WRS and S-WRS XRD patterns are available in Supplementary material (Fig. S1). Thus, rare earth diphenylphosphinates are found to have the same diffraction pattern, suggesting that [RE(dpp)]₃n are isostructural. The presence of an intense low-angle diffraction, close to seven 2θ/degrees (interplanar spacing d ~ 12 Å), is also repeated in all the synthesized coordination compounds. These results are similar to the behavior shown by MOFs [38] and corroborate those described in the literature [26,28,29,32].

However, the results revealed that a meticulous control of the reaction medium temperature and atmosphere was able to support the crystalline organization of the materials, significantly increasing the signal-to-noise ratio of the diffraction (Fig. 1).

When comparing interplanar distances of [Eu(dpp)]₃n reflections with those reported by Stucchi et al. (Table 1), small displacements in d (Å) values are observed, suggesting possible modifications in crystalline
The formation of [Eu(dpp)]\textsubscript{3} synthesized by the different routes, was also confirmed by displacements of \((POO-)\text{ass.}\) and \((POO-)\text{s.}\) stretching bands of the coordination compounds in relation to \((P = O)\) and \((PO(H)\) stretches of the ligand, respectively, which confirm that the oxygen of the phosphinate group has been coordinated to RE\textsuperscript{3+} (Table 2).

Coordination systematic was verified by comparing \((POO-)\text{ass.}\) and \((POO-)\text{s.}\) of [Eu(dpp)]\textsubscript{3} with respect to the same vibrational modes present in Na(dpp) and NH\textsubscript{4}(dpp) ionic compounds (Table 3). This determined coordination systematic follows trends similar to carboxylates ranges related to coordination modes. Thus, it may be considered that the oxygen of the phosphinate group has been coordinated to RE\textsuperscript{3+}.

Note: the p-value corresponding to the F-statistic of one-way ANOVA for \(d(\text{Å})\) values is higher than 0.05, suggesting that the treatments are not significant for that level of significance.

Regarding morphology, Francisco et al. [30] observed that [Y(dpp)]\textsubscript{3} particles obtained by those approaches can be found in Table 4. The high intensity sonication employed in U-WRS approach caused defined shape. The results of Rosa et al. [36] corroborate those described by Francisco et al. [30], whereby diphenylphosphinates complexes of La\textsuperscript{3+} -Ce\textsuperscript{3+} -Eu\textsuperscript{3+} / La\textsuperscript{3+} -Ce\textsuperscript{3+} -Tb\textsuperscript{3+} ternary systems prepared by precipitation are presented as agglomerates, with no defined shape. Through the synthetic approaches described in Subsections 2.1-2.4, excellent improvements in morphological characteristics of [Eu(dpp)]\textsubscript{3} were obtained, verified by SEM-FEG photomicrographs (Fig. 2).

Table 1

<table>
<thead>
<tr>
<th>WRS</th>
<th>U-WRS</th>
<th>M-WRS</th>
<th>S-WRS</th>
<th>Conventional WRS (provided by reference)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20 (degrees)</td>
<td>1/(I_0) (%)</td>
<td>(d(\text{Å}))</td>
<td>20 (degrees)</td>
<td>1/(I_0) (%)</td>
</tr>
<tr>
<td>7.28</td>
<td>100</td>
<td>12.14</td>
<td>7.28</td>
<td>100</td>
</tr>
<tr>
<td>14.64</td>
<td>7</td>
<td>6.05</td>
<td>14.64</td>
<td>7</td>
</tr>
<tr>
<td>17.62</td>
<td>5</td>
<td>5.03</td>
<td>17.60</td>
<td>7</td>
</tr>
<tr>
<td>19.44</td>
<td>29</td>
<td>4.57</td>
<td>19.44</td>
<td>32</td>
</tr>
<tr>
<td>20.50</td>
<td>15</td>
<td>4.33</td>
<td>20.48</td>
<td>19</td>
</tr>
<tr>
<td>22.98</td>
<td>9</td>
<td>3.87</td>
<td>23.04</td>
<td>10</td>
</tr>
<tr>
<td>24.14</td>
<td>6</td>
<td>3.69</td>
<td>24.10</td>
<td>7</td>
</tr>
<tr>
<td>25.26</td>
<td>10</td>
<td>3.53</td>
<td>25.24</td>
<td>12</td>
</tr>
</tbody>
</table>

Note: the p-value corresponding to the F-statistic of one-way ANOVA for \(d(\text{Å})\) values is higher than 0.05, suggesting that the treatments are not significant for that level of significance.

Table 3

<table>
<thead>
<tr>
<th>FTIR – (\Delta \nu(POO)) displacements of [Eu(dpp)]\textsubscript{3} and ionic compounds.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Synthetic route</td>
</tr>
<tr>
<td>-----------------</td>
</tr>
<tr>
<td>WRS</td>
</tr>
<tr>
<td>U-WRS</td>
</tr>
<tr>
<td>M-WRS</td>
</tr>
<tr>
<td>S-WRS</td>
</tr>
<tr>
<td>IC</td>
</tr>
<tr>
<td>NH\textsubscript{4}(dpp)</td>
</tr>
</tbody>
</table>

Units: cm\textsuperscript{-1}. Ref. [39-43].
damage to the crystalline structures and allowed the random breaking of the stick-shaped particles, decreasing their lengths by approximately 36% compared to those obtained via WRS, while preserving the diameters.

On the other hand, the use of the synthesis modulating agent provided a better kinetic control of nucleation. Thus, it allowed considerable elongation of the particles by reducing the crystal growth rate.

The solvothermal treatment, for providing dynamic conditions capable to dissolve and recrystallize materials that are relatively insoluble under ordinary conditions, was the approach that best managed to lengthen the 1D stick-shaped particles. In fact, particles 14 times longer than those obtained by WRS are observed, maintaining preserved their diameters in the range of 45–80 nm.

Solid state photoluminescence properties of WRS, U-WRS, M-WRS and S-WRS \([\text{Eu(dpp)}_3]_n\) were investigated by 77 K – PLS. All excitation (EX) spectra were obtained by monitoring \(\text{Eu}^{3+} 5D_{0} \rightarrow 7F_{1}\) transition, through which it is possible to observe (i) a broad and intense band related to \(^{1}A_{1g} \rightarrow ^{1}B_{2u}\) transitions of Ph(π-π*) system always around 270 nm and (ii) characteristic bands of Eu\(^{3+}\) between 310 and 500 nm referring to electronic transitions of the ground state \(^5F_0\) to various \(^{2S+1}L_J\) excited states, of which the bands close to 391 and 463 nm refer to Eu\(^{3+}\) 4f\(^6\) intra shell \(^7F_0 \rightarrow ^5L_{6}\) and \(^7F_0 \rightarrow ^5D_2\) transitions,

<table>
<thead>
<tr>
<th>Synthetic route</th>
<th>n length / nm</th>
<th>n diameter / nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>WRS</td>
<td>500 782 (± 499)</td>
<td>100 46.04 (± 1.08)</td>
</tr>
<tr>
<td>U-WRS</td>
<td>100 450 (± 197)</td>
<td>100 45.09 (± 0.07)</td>
</tr>
<tr>
<td>M-WRS</td>
<td>100 1,580 (± 730)</td>
<td>100 39.49 (± 0.30)</td>
</tr>
<tr>
<td>S-WRS</td>
<td>100 &gt; &gt; 1,000 (1,000 – 10,000)</td>
<td>100 63.10 (± 1.34)</td>
</tr>
</tbody>
</table>

Note: length and diameter are shown as the mean ± standard deviation, except for S-WRS particles length given the difficulty in counting intact particles due to their interlacing (see Fig. S3. - Supplementary material).

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**Fig. 2.** SEM-FEG photomicrographs @ 25,000 x of (a) WRS [Eu(dpp)_3]_n, (b) U-WRS [Eu(dpp)_3]_n, (c) M-WRS [Eu(dpp)_3]_n, (d) S-WRS [Eu(dpp)_3]_n. Note: other photomicrographs are available in Supplementary material (Fig. S3).

**Table 4**
Dimensions (length and diameter) of [Eu(dpp)_3]_n particles obtained by different synthetic routes.

**Fig. 3.** Statistical distribution of WRS [Eu(dpp)_3]_n particles dimensions: (a) length and (b) diameter.
Table 5

Integrated intensity ratio (R) $I(5D_{0} \rightarrow 7F_{2}) / I(5D_{0} \rightarrow 7F_{1})$.

<table>
<thead>
<tr>
<th>Synthetic route</th>
<th>$\lambda_{exc}$ (ligand)</th>
<th>$I_{0,2}/I_{0,1}$</th>
<th>$\lambda_{exc} (5F_{0} \rightarrow 7L_{6})$</th>
<th>$I_{0,2}/I_{0,1}$</th>
<th>$\lambda_{exc} (5F_{0} \rightarrow 7D_{2})$</th>
<th>$I_{0,2}/I_{0,1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>WRS</td>
<td>270.0</td>
<td>0.676 $^{\mathrm{a,b}}$ ± 0.070</td>
<td>391.4</td>
<td>0.621 $^{\mathrm{a,b}}$ ± 0.026</td>
<td>463.4</td>
<td>1.246 $^{\mathrm{a}}$ ± 0.023</td>
</tr>
<tr>
<td>U-WRS</td>
<td>268.4</td>
<td>0.564 $^{\mathrm{a,b}}$ ± 0.060</td>
<td>390.8</td>
<td>0.681 $^{\mathrm{a,b}}$ ± 0.069</td>
<td>462.8</td>
<td>1.064 $^{\mathrm{a}}$ ± 0.034</td>
</tr>
<tr>
<td>M-WRS</td>
<td>270.0</td>
<td>0.593 $^{\mathrm{a,b}}$ ± 0.051</td>
<td>391.8</td>
<td>0.640 $^{\mathrm{a,b}}$ ± 0.016</td>
<td>463.8</td>
<td>1.148 $^{\mathrm{a}}$ ± 0.059</td>
</tr>
<tr>
<td>S-WRS</td>
<td>269.0</td>
<td>0.428 $^{\mathrm{a,b}}$ ± 0.029</td>
<td>391.4</td>
<td>0.668 $^{\mathrm{a,b}}$ ± 0.055</td>
<td>463.4</td>
<td>0.840 $^{\mathrm{a}}$ ± 0.047</td>
</tr>
</tbody>
</table>

$\lambda_{exc}$ units: nm. $I_{0,2}/I_{0,1}$ are shown as the mean ± standard deviation. Different lowercase letters in the same row indicate statistically significant differences between the $\lambda_{exc}$ and different capital letters in the same column indicate statistically significant differences between the synthetic routes, according to Tukey’s test at 1% significance level. Other statistical data can be found in Supplementary material (Table S1 and Table S2).

respectively. From emission (EM) spectra it was possible to verify relaxations of the first Eu$^{3+}$ excited state, $5D_{0}$, to the first five levels of $7F_{J}$; it was also noted that the relative intensities of the emission spectrum bands have strong dependence on the excitation wavelength ($\lambda_{exc}$) and that these bands undergo influence of the aromatic ring-phosphoryl conjugate system excitation band up to 421 nm. Bands from this wavelength could be basically assigned to Eu$^{3+}$.

Eu$^{3+}$ $5D_{0} \rightarrow 7F_{1}$ (0–1) transition follows magnetic dipole rules, which makes it extremely independent of Eu$^{3+}$ chemical environment [46]. However, the intensity of this transition can be identified to be constant if considered its total integrated intensity. On the other hand, Eu$^{3+}$ $5D_{0} \rightarrow 7F_{2}$ (0–2) transition, governed by electric dipole mechanism, is known to be hypersensitive to the chemical environment occupied by Eu$^{3+}$ and the nature of the ligands; its intensity is proportional to Judd-Ofelt $\Omega_2$ parameter [47].

Whereas there are no modifications in ligand compositions and no substantial modifications at the metal-ligand bonding distances, we can use the ratio (R) between the integrated intensities of 0–2 and 0–1 transitions as evidence of the existence of more than one type of symmetry site occupied by Eu$^{3+}$ (Table 5). However, it is important to clarify that the ratio R is dependent on factors beyond the point symmetry of Eu$^{3+}$ ion, such as the coordination polyhedron structure [48].

The EM and EX spectra for WRS [Eu(dpp)$_3$], in Fig. 4. In the EM spectrum (Fig. 4(b)), we found two distinct situations: (i) when the excitation is centered on the ligand $^1A_{1g} \rightarrow ^1B_{2u}$ transitions of Ph (π–π$^*$) system – or the Eu$^{3+}$ $7F_0 \rightarrow 5L_6$ transition, there is predilection for symmetric sites over the lower symmetry ones, i.e. the symmetric sites contribute preferentially to the Eu$^{3+}$ emission and (ii) the excitation in Eu$^{3+}$ $7F_0 \rightarrow 5D_2$ intraconfigurational transition increases the contribution of lower symmetry sites for the intensity of EM spectrum bands. It is important to emphasize that the Eu$^{3+}$ $7F_0 \rightarrow 5L_6$ transition remains sensitive to the electronic transition of the ligand whereas when the excitation is centered on Eu$^{3+}$ $7F_0 \rightarrow 5D_2$ transition that influence is practically eliminated.

U-WRS, M-WRS and S-WRS methodologies provided a homogeneous nucleation and a controlled recrystallization rate for [Eu(dpp)$_3$]$\text{Na}$ preparations, leading to a significant increase in the crystallinity of the material and an improvement in its morphological quality. Thereby, changes in precipitation kinetics and crystalline phases formation rate generated a distribution of Eu$^{3+}$ ions in higher symmetry sites (HSS), so that the contribution of the lower symmetry sites never became equal to or greater than that observed in WRS methodology.

S-WRS provided a better ordering of [Eu(dpp)$_3$]$\text{Na}$ structure by facilitating the interaction between the reactional components, achieving thermodynamically stable and metastable states and thus contributing to the formation of new materials which could not be easily synthesized from conventional synthetic routes. This synthetic approach made it possible to significantly reduce the contribution of the less symmetrical site (LSS). In fact, this LSS contribution to Eu$^{3+}$ emission never becomes preponderant, not even when the excitation at Eu$^{3+}$ $7F_2 \rightarrow 5D_2$ transition (Fig. 5).

It is known that previous studies[28,29,32] have already pointed to the possibility of Eu$^{3+}$ occupying more than one symmetry site in yttrium, lanthanum and gadolinium diphenylphosphinates matrices doped with trivalent europium. However, it is the first time that the existence of at least two symmetry sites is verified in a pure europium diphenylphosphinate matrix [Eu(dpp)$_3$]$\text{Na}$ synthesized by different methodologies.

4. Conclusions

The new methodologies proposed in this work for [Eu(dpp)$_3$]$\text{Na}$ synthesis rendered crystalline and organized materials as a result of meticulous control of the synthesis parameters. Actually, such strict control led to the formation of 1D nanostructured materials in stick-shaped particles. In addition, comparisons of v(POO)$_\text{ass}$ and v(POO)$_\text{k}$ stretches present in [Eu(dpp)$_3$]$\text{Na}$, Na(dpp) and NH$_4$(dpp) made it possible to determine that [Eu(dpp)$_3$]$\text{Na}$ are compounds in which chelate and bridged coordination modes coexist. The relative intensities of EM spectrum bands have strong dependence on the excitation wavelength. In an unprecedented way, we could observe, through the integrated intensity ratio (R) $I(5D_0 \rightarrow 7F_2) / I(5D_0 \rightarrow 7F_1)$, the existence of more than one symmetry site type occupied by Eu$^{3+}$ in a pure europium diphenylphosphinate matrix. We were also able to show that U-WRS,
M-WRS and S-WRS methodologies provided a distribution of Eu$^{3+}$ ions in higher symmetry sites (HSS); the contribution of the sites of lower symmetry to the emission never became equal to or greater than that observed in WRS. As a matter of fact, S-WRS was the approach that best ordered Eu(dpp)$_3$, structure, reducing significantly the contribution of the less symmetrical site (LSS).

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at https://doi.org/10.1016/j.jlumin.2018.06.040.

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