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$La_2(CN_2)_3$ – the missing link of rare-earth carbodiimides, prepared through an efficient synthetic route and its Ce^{3+} activated photoluminescence;

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Rare-earth (RE) carbodiimides according to the composition $RE_2(CN_2)_3$ have been reported for the whole series of RE elements, all prepared by solid-state metathesis (SSM) reactions. Only one compound, $La_2(CN_2)_3$, could not be made by this way of synthesis. Herein, we report the preparation of $La_2(CN_2)_3$ by using lanthanum cyanurate as a single-source precursor. The conversion of the precursor is analyzed by thermoanalytical studies. The crystal structure of the precursor and the novel $La_2(CN_2)_3$ are characterized by X-ray diffraction techniques. $La_2(CN_2)_3$ is represented by a distinct crystal structure with a dodecahedral environment of the La^{3+} ion. Having the knowledge of the last missing rare-earth carbodiimide, we herein present a summary of all existing $RE_2(CN_2)_3$ compounds, including their structural relationships. Doping with Ce^{3+} leads to the $La_2(CN_2)_3$: Ce^{3+} phosphor, which is reported with its photoluminescence properties.

Introduction

Metal dinitridocarbonates, commonly denoted as metal cyanamides (N=C-N²-) and metal carbodiimides (¬N=C=N¬), have been reported by different ways of synthesis. The first method for synthesizing a carbodiimide involves reacting calcium carbide (CaC₂) with nitrogen gas (N₂) at elevated temperatures, approximately 1000 °C, in a process known as the Frank-Caro method. 1

Following the discovery of calcium carbodiimide, numerous metal carbodiimide or metal cyanamide compounds were developed. An early synthetic approach to lithium carbodiimide ($\text{Li}_2(\text{CN}_2)$) was achieved in the 1970s by heating Li_3N and Li_2C_2 together at about 600 °C.² Another proposed method for the synthesis of $\text{Li}_2(\text{CN}_2)$ is ammonolysis, which involves the reaction of lithium carbonates (Li_2CO_3) with ammonia (NH₃).³ Alternatively, lithium carbodiimide ($\text{Li}_2(\text{CN}_2)$) can be synthesized using lithium nitride (Li_3N)⁴ (eqn (1)) or hydride (LiH)⁵ and melamine. For synthesizing other alkali metal carbodiimides such as sodium⁴ and potass-

ium⁷ carbodiimides an alternative route was employed. Alkaline-earth metal carbodiimides such as Mg(CN₂), Sr(CN₂), and Ba(CN₂) can be produced similarly utilizing the reaction of alkaline-earth metal nitrides with melamine at temperatures ranging from 740 to 850 °C.⁸

One of the most effective approaches for synthesizing many other metal carbodiimides is the solid-state metathesis (SSM), which allows for the production of relatively pure carbodiimides under moderate heating conditions, typically at 450-600 °C. ^{9,10} Within this process, a metal halide (MX₂ or MX₃) is converted with lithium carbodiimide in a salt-balanced reaction (eqn (2)) to yield a metal carbodiimide.

$$2Li_3N + C_3H_6N_6 \rightarrow 3Li_2(CN_2) + 2NH_3$$
 (1)

$$2MX_3 + 3Li_2(CN_2) \rightarrow M_2(CN_2)_3 + 6LiX$$
 (2)

To produce calcium carbodiimide of high purity, another synthetic route was developed as shown in eqn (3). This methodology was first proposed by Seifer¹¹ and then further refined in 2023.¹² The study done by Seifer suggests briefly the preparation of lead, barium, and strontium carbodiimide in addition to calcium carbodiimide. These carbodiimides are prepared by the reaction of cyanuric acid with metal chlorides in the presence of sodium hydroxide in water to produce the corresponding cyanurate salts which are subsequently pyrolyzed to give the desired carbodiimides. This synthetic route shows the promising route of using triazine-derived precur-

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sors, for example, melamine,^{5,13} and derivatives of cyanuric acid in the production of carbodiimides.

$$Ca(HO_3C_3N_3) \rightarrow Ca(CN_2) + HOCN + CO_2 \uparrow$$
 (3)

It is noteworthy to mention that some transition metal carbodiimides were also synthesized using aqueous solution-based methods for instance, $Zn(CN_2)$, 14 $Co(CN_2)$, 15 $Ni(CN_2)$, 15 $Cu(CN_2)$, 16 $Cd(CN_2)$, 17 $Ag_2(CN_2)$, 18 $Hg(CN_2)$. 19 Other transition metal carbodiimides, such as $Mn(CN_2)$, 20 $Cr_2(CN_2)$, 21 Zr (CN_2) , 22 and $Hf(CN_2)$, 22 can be synthesized using the similar solid-state metathesis (SSM) reactions between a metal halide with $Li_2(CN_2)$ or $Zn(CN_2)$, 14 p-Block metal carbodiimides, like $Pb(CN_2)$, 23 $Bi_2(CN_2)$, 24 $Tl_2(CN_2)$, 25 and $In_{2.24}(CN_2)$, 13,26 are synthesized by reacting metal salts with cyanamide or cyanide compounds. For tin carbodiimides, the conventional SSM reaction of $Li_2(CN_2)$ with metal halides such as $SnCl_2$ or SnF_2 , yields $Sn(CN_2)$ and $Sn_4Cl_2(CN_2)$ 3. $Sn^{3,27}$

SSM reactions have been extensively utilized by our research group to synthesize rare-earth (RE) carbodiimides, essentially by reacting lithium carbodiimide with a rare-earth metal chloride. 28,29 Most prominent is the series of RE2(CN2)3 compounds that have been reported for RE elements from Sc to Lu, except for RE = La (and the radioactive element Pm). $^{28-30}$ Lanthanum carbodiimide was first reported 76 years ago from reactions of La₂O₃ with HCN.³¹ However, the products were poorly characterized, with no structural or spectroscopic data available.31 It is worth mentioning that compounds of smaller rare-earth ions like RE = Sc,³² Tm, Yb, and Lu^{28,33} crystallize with the trigonal rhombohedral space group $R\bar{3}c$ (Z = 6), and the coordination number (CN) six of the RE³⁺ ion.²⁸ Corresponding compounds with RE = Y and Ce-Er (except Pm) crystallize monoclinically with the space group C2/m (Z = 2) and the CN of the RE³⁺ being seven. ^{28,33,34} Tm₂(CN₂)₃ has been shown to undergo a pressure transformation from $R\bar{3}c$ into C2/m with a significant volume reduction.²⁸

Through this route, not only pseudobinary but also pseudoternary, rare-earth (RE) compounds have been synthesized. These compounds cover a wide range of mixed cation and mixed anion carbodiimides such as RE₂O₂(CN₂) (RE = Ce, Pr, Nd, Sm, Eu, Gd, Dy-Yb), 35,36 RECl(CN₂) (RE = La-Pr), 37 Sc₂O (CN_2) , ³² RE₂Cl (CN_2) N (RE = La, Ce), ³⁸ RE₂Br (CN_2) N (RE = La, Pr), ³⁹ $REI(CN_2)N$ (RE = La, Gd), ⁴⁰ and $Eu_2I_2(CN_2)$, ⁴¹ Eu₄F₅(CN₂)₂.⁴² Furthermore, some pseudoquaternary NCN rare-earth (RE) compounds containing three different cations have been also developed by adding the third reactant to the conventional SSM reactions. Examples of these compounds carbidonitridosilicates,34,43,44 rare-earth tetracyanamidogermanates. 45,46 For lanthanum, mixed anion compounds $La_2O_2(CN_2)^{47}$ $La_2S_2(CN_2)^{48}$ $La_2O(CN_2)^{37}$ La₃(CN₂)₃N⁴⁹ or LaCl(CN₂)⁵⁰ have been so far reported.

The photoluminescence properties of lanthanide (Ln) doped $RE_2(CN_2)_3^{29}$ such as $Gd_2(CN_2)_3$:Ce or Tb, Ce, and Tb²⁹ have been thoroughly studied, leading to the development of a pc-LED prototype-based on $Y_2(CN_2)_3$:Ce.⁵¹ In Ce³⁺ doped materials luminescence usually occurs due to electronic tran-

sition from ground state levels of $(^2F_{5/2}$ and $^2F_{7/2})$ of the [Xe]4f¹ configuration to lowest crystal-field components of the [Xe]5d¹. By tuning the crystal-field strength and covalent character of Ce³⁺, the material's luminescence properties can be adjusted.²⁹

Herein we explore the preparation of lanthanum carbodiimides through this efficient way of synthesis. We describe the formation and structural characterization of intermediate lanthanum cyanurates obtained from aqueous solution and the thermal conversion into the novel $La_2(CN_2)_3$. The $La_2(CN_2)_3$ represents the missing compound among the series of $RE_2(CN_2)_3$ compounds with a new crystal structure. The dodecahedral coordination of the lanthanum ion in $La_2(CN_2)_3$ parallels the coordination pattern of yttrium in the structure of yttrium aluminum garnet (YAG), which resembles the host structure for the most prominent YAG:Ce phosphor in todaýs phosphor converted light-emitting diodes (pc-LED).

Results and discussion

Preparation

Rare-earth carbodiimides were successfully prepared by means of solid-state metathesis reaction (see eqn (2)). However, this way of synthesis has been unsuccessful for $La_2(CN_2)_3$, because all reactions have led to the formation of $LaCl(CN_2)_3^{37}$

Herein, we report the preparation of $\text{La}_2(\text{CN}_2)_3$ *via* a precursor route. The compound was synthesized through the thermal decomposition of a lanthanum cyanurate precursor. This precursor was obtained by reacting lanthanum chloride with cyanuric acid and sodium hydroxide in an aqueous solution, yielding an insoluble precipitate eqn (4).

$$\begin{aligned} & LaCl_{3} + 2H_{3}C_{3}N_{3}O_{3} + 3NaOH \\ & \rightarrow La(H_{2}O_{3}C_{3}N_{3})(HO_{3}C_{3}N_{3})(H_{2}O)_{3} \downarrow + 3NaCl \end{aligned} \tag{4}$$

It is noteworthy that two distinct phases of lanthanum cyanurates were obtained as insoluble precipitates. We found out that the formation of these two phases was dependent on the pH of the solution which can be controlled by varying the amount of water content. Lower amounts of water and thus higher pH values led to the formation of a lanthanum cyanurate composed of a single deprotonated and a double deprotonated cyanurate (La(HC₃N₃O₃)(H₂C₃N₃O₃)(H₂O)₃). In contrast, higher amounts of water and thus lower pH values led to the formation of a lanthanum cyanurate with two single deprotonated cyanurates along with a hydroxide ion (La $(H_2C_3N_3O_3)_2(OH)(H_2O)_4\cdot H_2O)$. These results can be explained by the pK_a values of the first two deprotonation steps of cyanuric acid (p K_{a1} = 6.88 and p K_{a2} = 11.40).⁵² Only at higher pH values, the solution becomes sufficiently basic to allow the second deprotonation of the cyanuric acid and thus the formation of the lanthanum cyanurate, whereby the cyanuric acid unit is present in its double deprotonated form. However, thermoanalytical studies in the next section, will demonstrate that both lanthanum cyanurate phases could be converted to

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lanthanum carbodiimide at 770 °C under a flow of argon. The lanthanum carbodiimide demonstrated air and water stability over a period of four weeks, as confirmed by X-ray powder diffraction.

Thermoanalytic studies

The thermal decomposition of lanthanum cyanurates was investigated using thermal analysis methods such as thermogravimetric analysis (TGA) combined with differential thermal analysis (DTA). The measurements were made in an argon flow within a temperature range from room temperature up to 770 °C at a rate of 2 K min⁻¹ for both heating and cooling. Fig. 1 represents the TGA of La(HC₃N₃O₃)(H₂C₃N₃O₃)(H₂O)₃ which shows that this material undergoes a decomposition in three steps, ultimately leading to the formation of lanthanum carbodiimide. Based on the measured and theoretical mass losses at each stage, the first step at 187 °C, with a mass loss of -12.04% (theory: -12.15%), is attributed to the loss of water (eqn (5)). The second step at 405 °C, with a mass loss of -28.38% (theoretical: -29.03%), can be corresponded to the decomposition of La(HC₃N₃O₃)(H₂C₃N₃O₃) to La(C₃N₃O₃) which is a completely deprotonated unit (eqn (6)). The final step, occurring at 737 °C with a mass loss of -14.82% (theory: -14.85%), is associated with the release of carbon dioxide, resulting in the formation of lanthanum carbodiimide, La₂(CN₂)₃ (eqn (7)). Heating the lanthanum carbodiimide to even higher temperatures (1000 °C), resulted in the formation of lanthanum nitride (LaN). This sequence can be summarized in the following reactions:

$$2La(HC_3N_3O_3)(H_2C_3N_3O_3)(H_2O)_3$$

$$\to 2La(HC_3N_3O_3)(H_2C_3N_3O_3) + 6H_2O \uparrow$$
 (5)

$$2La(HC_3N_3O_3)(H_2C_3N_3O_3) \to 2La(C_3N_3O_3) + 2H_3C_3N_3O_3 \uparrow$$
(6)

$$2La(C_3N_3O_3) \rightarrow La_2(CN_2)_3 + 3CO_2 \uparrow \tag{7}$$

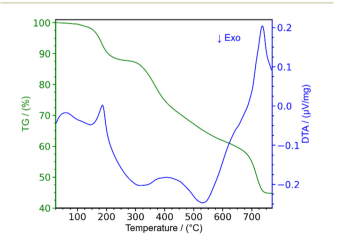


Fig. 1 Thermogravimetric analysis (TGA) combined with differential thermal analysis (DTA) of La(HC₃N₃O₃)(H₂C₃N₃O₃)(H₂O)₃.

Crystal structures

The synthesis of La₂(CN₂)₃ can be achieved by the decomposition of a single source precursor, which can be either $La(HC_3N_3O_3)(H_2C_3N_3O_3)(H_2O)_3$ (1) or $La(H_2C_3N_3O_3)_2(OH)$ (H₂O)₄·H₂O (2). Crystal structures of both precursors were refined based on single-crystal X-ray diffraction, with relevant data summarized in Table 1. Both compounds contain one type of La³⁺ in the structure. The first compound (1) contains mono- and divalent cvanurate anions and three water molecules in the coordination environment of the La3+, the latter (2) monovalent cyanurate anions, bridging hydroxide and four water molecules as shown in Fig. 2 and 3. There are at least two more lanthanum cyanurate structures being reported in the literature. The crystal structure of La[H₂N₃C₃O₃]₃·8.5 H₂O ⁵³ was refined with the space group P1, containing monovalent cyanurate anions. Another structure, described as La (H₂C₃N₃O₃)₂·OH·2H₂O, ⁵⁴ leaves a precise assignment of hydrogen atoms behind.

The appearance of a number of different lanthanum cyanurate compounds emphasizes the influence of pH conditions during precipitation in aqueous solution and by a varying extent of hydration. However, so far there is no indication that the decomposition of any of these precursors would not lead to $La_2(CN_2)_3$.

The crystal structure of $La(HC_3N_3O_3)(H_2C_3N_3O_3)(H_2O)_3$ (1) is characterized by one type of lanthanum, being surrounded by three coordinated water molecules and five cyanurate ions showing a bridging functionality (Fig. 2).

The crystal structure of La₂(CN₂)₃ was solved and refined on the basis of X-ray powder diffraction data by Rietveld refine-

Table 1 Crystallographic data of cyanurate precursor compounds La $(HC_3N_3O_3)(H_2C_3N_3O_3)(H_2O)_3$ (1), $La(H_2C_3N_3O_3)_2(OH)(H_2O)_4\cdot H_2O$ (2) and of La2(CN2)3

	(1)	(2)	$La_2(CN_2)_3$
CCDC code	2403637	2409323	2393819
Empirical formula	C ₆ H ₉ LaN ₆ O ₉	$C_6H_{15}LaN_6O_{12}$	$C_3La_2N_6$
Formula weight (g mol ⁻¹)	448.10	502.15	397.88
Crystal system	Monoclinic	Triclinic	Monoclinic
Space group	$P2_1/c$	$P\bar{1}$	I2/a
a/Å	8.0793(4)	6.3215(3)	8.69003(6)
b/Å	17.1808(7)	11.0993(6)	6.88968(5)
c/Å	8.5107(4)	11.8544(5)	10.30517(9)
<i>α</i> /°		66.312(5)	
β/°	99.223(5)	88.178(3)	105.0711(6)
γ/°		75.768(4)	
$V/\text{Å}^3$	1166.09(9)	736.20(7)	595.764(8)
Z	4	2	4
μ/mm^{-1}	29.027	23.237	108.706
$D_{\rm c}/{\rm g~cm^{-3}}$	2.552	2.265	4.436
Crystal size	$0.05\times0.03\times0.01$	$0.04\times0.04\times0.01$	Powder
Θ range/°	5.149 to 74.464	4.084 to 72.102	2.5 to 60
Reflections collected	41 457	22 199	488
Parameters	235	251	45
R_{Bragg}	_	_	4.0955
χ^2	_	_	1.0244
Wavelength (Cu- K_{α}) (Å)	1.54184	1.54184	1.54184
R_1 , w R_2 $(I > 2\sigma(I))$	0.0162, 0.0385	0.0333, 0.0837	_
R indices (all data)	0.0175, 0.0389	0.0360, 0.0854	_
GOOF	1.041	1.076	1.012

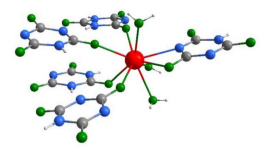


Fig. 2 Coordination environment of the La^{3+} in the structure of $La(HC_3N_3O_3)(H_2C_3N_3O_3)(H_2O)_3$ (1) (gray: C, blue: N, green: O, red: La, white: H).

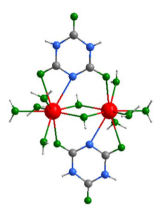


Fig. 3 Coordination environment of the La^{3+} ion in the structure of La $(H_2C_3N_3O_3)_2(OH)(H_2O)_4\cdot H_2O$ (2) (gray: C, blue: N, green: O, red: La, white: H).

ment with the space group I^2/a (Table 1 and Fig. 4). Crystal structures of RE carbodiimides follow a characteristic structure pattern in which the $[N=C=N]^{2-}$ ions are arranged in layers, following the motif of a hexagonal closed packing of sticks. Cations are situated in between these layers to form an alternating arrangement.

This pattern is apparent also in the structure of $La_2(CN_2)_3$, displayed in Fig. 5. Major differences among the structures of $RE_2(CN_2)_3$ compounds are the tilting of N=C=N ions within layers relative to each other, and the exact position of RE ions in structures. The crystal structure of $La_2(CN_2)_3$ contains one type of lanthanum in the structure and two distinct carbodiimides ions. Lanthanum ions in the structure have the coordination number eight, shown in Fig. 6, together with the pattern of the corresponding dodecahedron.

The characterization of $\text{La}_2(\text{CN}_2)_3$ completes the series of binary rare-earth carbodiimides, which is represented by three distinct structures, respectively, coordination patterns (Fig. 7), with their unit volumes displayed in Fig. 8. The general trend of molar unit cell volumes represents the lanthanide contraction for the series from Ce to Tm with the space group C2/m (Z=2) with the coordination number (CN) of the RE³⁺ being seven. Compounds of Tm, Yb, and Lu follow the same trend,

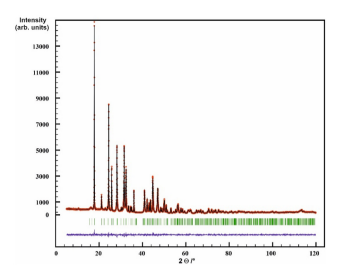


Fig. 4 Rietveld refinement of the XRD powder pattern of $La_2(CN_2)_3$. Observed intensities are marked as red circles, and calculated intensities as black lines. Bragg positions are marked as green lines, the difference curve $I_{observed} - I_{calculated}$ as blue line.

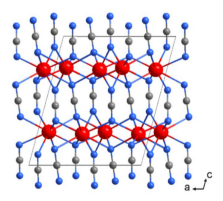


Fig. 5 The layered appearance of the crystal structure of $La_2(CN_2)_3$ with lanthanum ions is shown in red.

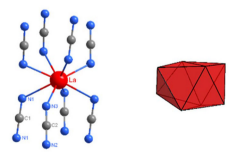


Fig. 6 The environment of lanthanum with two crystallographically distinct $[N=C=N]^{2-}$ ions (distances: C1–N1 = 124.41(6) pm, \angle N1–C1–N1 = 177.82(4)°; C2–N2 = 123.95(9) pm, C2–N3 = 124.17(9) pm, \angle N2–C2–N3 = 178.55(7)°) and a dodecahedron representing the coordination environment of La $^{3+}$ in the structure of La 2 (CN $_{2}$) $_{3}$.

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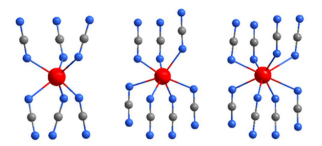


Fig. 7 Coordination environments of rare-earth ions (red) in the trigonal $(R\bar{3}c)$, monoclinic (C2/m), and (I2/a) structures, from left to right.

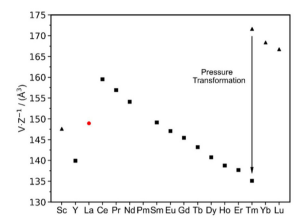


Fig. 8 Molar volume (V/Z) of rare-earth carbodiimides, RE₂(CN₂)₃, with the monoclinic (C2/m) structure for the series RE = Y, and Ce-Tm (displayed as squares), the trigonal $(R\bar{3}c)$ structure for RE = Sc (displayed as triangles) and Tm-Lu, and monoclinic (12/a) La₂(CN₂)₃ (displayed as a red dot). $Tm_2(CN_2)_3$ has been shown to be dimorphic with the monoclinic (C2/m) structure and the trigonal $(R\overline{3}c)$ structure of the highpressure phase.

however with a trigonal structure with the CN of the RE being six. Tm₂(CN₂)₃ is dimorphic and undergoes a pressure transformation from $R\bar{3}c$ into C2/m. Thereby, the coordination number of the Tm³⁺ ion increases from six to seven. Lanthanum, as the largest lanthanide ion, appears with the coordination number eight.

The lanthanum site with its dodecahedral environment in La₂(CN₂)₃ appears to be an interesting site for doping with a photoluminescence activator. A most prominent example of such a dodecahedral environment is apparent in the pc-LED phosphor Y₃Al₅O₁₂:Ce. Consequently, we conducted photoluminescence studies on La₂(CN₂)₃:Ce, discussed later in this work.

Infrared spectroscopy

The infrared (IR) spectrum of La₂(CN₂)₃ has also been reported to support its carbodiimide character (see Fig. 6 for distances and angles). As can be observed in Fig. 9 the characteristic vibrational bands of the [NCN]²⁻ ion at 2029 and 1929 cm⁻¹ can be ascribed to the asymmetric stretching vibrations (ν_{as}).

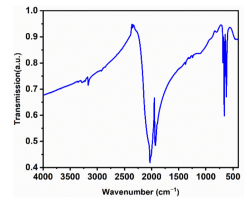


Fig. 9 FT-IR spectrum of La₂(CN₂)₃.

Furthermore, the corresponding bending vibrations (δ) are observed at 685, 660, and 621 cm⁻¹. The absence of a symmetric vibration (ν_s) in the IR spectrum supports the identification of the compound as a carbodiimide rather than a cyanamide.³⁷ Furthermore, the results align well with previously reported data on Table 2 for two other rare-earth metal carbodiimide modifications.55

Photoluminescence spectroscopy

The photoluminescence (PL) spectrum of La₂(CN₂)₃ doped with Ce3+ (5 atom% w.r.t La) has been studied to evaluate the potential of this novel material for application in a luminescent screen. Ce3+ is already well-known as an efficient emitter in various applications of luminescent materials. 56-59 For instance, Ce3+ is one of the most important emitters in LED phosphors, such as $Ln_3(Al,Ga,Sc)_5O_{12}$:Ce (Ln = Y, Gd, Tb, Lu), or acts as a primary activator ion in scintillators, including Lu₂SiO₅:Ce, LuAlO₃:Ce, and Lu₃Al₅O₁₂:Ce (LuAG:Ce³⁺) (see Table 3). Ce3+-activated materials also play an important role

Table 2 Vibrational frequencies (cm⁻¹) from lanthanum and selected rare-earth carbodiimides⁵⁵

	$\nu_{ m as}$ (CN ₂	$\nu_{\rm as} \left({\rm CN}_2\right)^{2-}$)2-		
$Lu_2(CN_2)_3$ $Sm_2(CN_2)_3$ $La_2(CN_2)_3$	2080 2023 2029	2009 1955 1929	— 705 685	680 668 660	640 634 621	616 —

Table 3 Emission maxima, density, and decay time of some Ce³⁺ activated phosphors and scintillators 61,62

Ce ³⁺ phosphor or scintillator	Emission max. nm	Density g cm ⁻³	Decay time
LaBr ₃ :Ce	358	5.3	35
YAlO ₃ :Ce	360	5.6	20-30
LuAlO ₃ :Ce	365	8.3	18
Lu ₂ SiO ₅ :Ce	390	7.4	30
Gd ₂ SiO ₅ :Ce	420	6.7	60
Lu ₃ Al ₅ O ₁₂ :Ce	525	6.7	54

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in UV lamps, such as YPO4:Ce, LaPO4:Ce, LaMgAl11O19:Ce, and YMgB₅O₁₀:Ce.⁶⁰

The photoluminescence of materials doped with Ce³⁺ ions originates from interconfigurational transitions, i.e. of transitions between different electronic configurations. In the ground state, Ce³⁺ has the configuration [Xe]4f¹ which, upon excitation is promoted to the configuration [Xe]5d1. The energy required for this transition depends on the crystal field splitting of the 5d levels, which is influenced by the surrounding coordinated anions and centroid shift caused by the local environment. The emission process caused by the relaxation of the 5d¹ to the 4f¹ configuration yields two broad emission bands due to the ground state term spin-orbit splitting yielding the terms ${}^{2}F_{5/2}$ and ${}^{2}F_{7/2}$. This kind of transition is spin

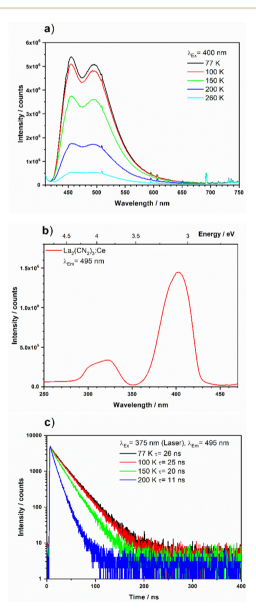


Fig. 10 (a) Emission spectra at several temperatures and (b) excitation spectra, and (c) decay curve upon 375 nm excitation of La₂(CN₂)₃: $Ce^{3+}(5\%)$

and parity allowed, therefore, resulting in a high oscillator strength and a short decay constant. Since a 5d orbital is involved, cerium-activated materials may show UV, blue, green, or yellow emission depending on the crystal-field-determined energetic position of absorption and emission bands. The 5d orbitals are more spatially extended and thus strongly interact with the crystal field around the atom, therefore, splitting into different levels.

Fig. 10a shows the broad emission band of La₂(CN₂)₃ doped with 5% of Ce³⁺, with two maxima at 450 nm (2222 cm⁻¹) and 500 nm (20 000 cm⁻¹) upon 400 nm excitation, at various temperatures. The excitation spectrum monitored for 495 nm is also shown in Fig. 10b and exhibits a strong excitation band at 400 nm (25 000 cm⁻¹), thus the Stokes shift is just 2800 cm⁻¹, which points to little relaxation in the excited state. As mentioned above, the crystal field strength and thus chemical environment such as ligand type, symmetry, and metal-toligand distance determines the energy gap between the [Xe]4f¹ and [Xe]5d¹ configuration. Therefore, the PL spectra will change if Ce³⁺ is located onto different crystallographic sites. A weaker crystal field increases the energy gap and results in a shift of the PL spectra towards higher energy such as UV or blue light. A comparison with the earlier published Gd₂(CN₂)₃: Ce³⁺ with a broad emission band at 575 nm (17 391 cm⁻¹) under 415 nm excitation,²⁹ and Y₂(CN₂)₃:Ce³⁺⁵¹ with a emission band range of 570-577 nm under 415 nm excitation, confirms that the emission bands in La₂(CN₂)₃:Ce³⁺ are strongly blue-shifted. This can be explained by the weaker crystal field in La2(CN2)3 than Gd2(CN2)3, and Y2(CN2)3 due to the larger metal-to-ligand distances onto the La3+ site. The material shows at low temperature a single exponential decay curve, while the calculated decay time of 26 ns at 77 K (Fig. 10c) is typical for blue-emitting Ce3+ activated luminescent materials or scintillators.

Conclusions

The vast majority of rare-earth carbodiimide compounds have been prepared by solid-state metathesis (SSM) reactions. However, the preparation of La₂(CN₂)₃ has yet failed by SSM, because its formation is hindered in favour of the formation of La₂Cl(CN₂). An alternative way of preparation of metal carbodiimides is the employment of metal cyanurate precursors, showing the formation of pure La₂(CN₂)₃ by decomposition of lanthanum cyanurate hydrate as a single-source precursor.

Lanthanum cyanurate compounds can appear with monoor divalent cyanurate anions and with different amounts of water molecules. The thermal conversion into lanthanum carbodiimide appears in three basic steps, by release of water, the loss of cyanuric acid, and finally the loss of CO₂.

The crystal structure of lanthanum carbodiimide completes the series of rare-earth carbodiimides that is represented by three distinct structure types and coordination numbers of the rare-earth ions. The structure of La₂(CN₂)₃ contains La³⁺ with the coordination number eight and represents an attractive

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host lattice for doping with rare-earth activators. Doping with Ce^{3+} leads to the luminescent material $La_2(CN_2)_3$:Ce which shows a blue photoluminescence on excitation with 375 nm. Due to its high density and short decay time, this phosphor can be considered as a perspective szintillator material.

Experimental section

Materials and methods

The following starting materials cyanuric acid (Sigma-Aldrich, 98%), lanthanum(III) chloride heptahydrate (Fluka AG, 99.9%), sodium hydroxide (Chemsolute, 98.8%), and cerium(III) chloride heptahydrate (Merck, 98.5%) were used without further purification and were handled under air and standard conditions.

The thermal decomposition was carried out in a Carbolite HST 12/300 furnace equipped with a 1 m long quartz glass furnace tube. The decomposition product was handled under an inert argon atmosphere and transferred into a glovebox with maintained moisture and oxygen levels below 1 ppm.

Synthesis

La(HC₃N₃O₃)(H₂C₃N₃O₃)(H₂O)₃ (1). 69.5 mg (538.6 mmol) cyanuric acid and 100.0 mg (269.3 mmol) lanthanum(III) chloride heptahydrate were combined in a beaker equipped with a magnetic stirrer. 10 mL deionized water was then introduced at room temperature and the mixture was subsequently heated to 100 °C. Upon complete dissolution of the cyanuric acid, 32.3 mg (807.8 mmol) sodium hydroxide was added to the reaction. The mixture was then stirred for 1 h.

The resulting precipitate was collected by filtration (yield w.r.t. La 64%), washed three times with 20 mL of deionized water to remove sodium chloride and dried in an oven at $87.5~^{\circ}\text{C}$.

La(H₂C₃N₃O₃)₂(OH)(H₂O)₄·H₂O (2). 69.5 mg (538.6 mmol) cyanuric acid and 32.3 mg (807.8 mmol) sodium hydroxide were combined in a beaker equipped with a magnetic stirrer. Then 30 mL deionized water was introduced at room temperature and the mixture was subsequently heated to 100 °C. Upon complete dissolution of the cyanuric acid, 100.0 mg (269.3 mmol) lanthanum(III) chloride heptahydrate in 10 mL deionized water was added dropwise to the reaction. The mixture was then stirred for 3 h. The resulting precipitate was collected by filtration (yield w.r.t. La 27%), washed three times with 20 mL of deionized water and dried in an oven at 87.5 °C.

La₂(CN₂)₃. 100 mg (223.2 mmol) lanthanum cyanurate (2) was placed in a tube furnace and then heat-treated under flowing Ar at 770 °C for 30 min with a ramping rate of 2 K min⁻¹. Following the holding time, the furnace was cooled to room temperature with the same ramping rate of 2 K min⁻¹, yielding the final product (yield w.r.t. La >99%).

 $\text{La}_2(\text{CN}_2)_3$:Ce. The synthesis for the doped lanthanum carbodiimide was performed similarly to the already established synthesis route. Hereby 200.0 mg (446.4 mmol) La $(\text{H}_2\text{O}_3\text{C}_3\text{N}_3)/(\text{HO}_3\text{C}_3\text{N}_3)/(\text{H}_2\text{O})_3$ was mixed and subsequently

mortared with 8.3 mg (22.3 mmol) cerium(III) chloride heptahydrate (CeCl₃·7H₂O) and then placed into the oven.

Thermoanalytic studies

Thermogravimetric analysis and differential thermal analysis were performed using a Netzsch Jupiter STA 449 F3 thermal analyzer. The synthesized samples were measured in an open corundum crucible under a continuous argon flow.

X-ray powder diffraction

Compounds were investigated by powder X-ray diffraction (XRD) on a Stadi-P (STOE, Darmstadt) diffractometer with germanium monochromated $\text{Cu-K}_{\alpha 1}$ radiation. The powder XRD pattern of $\text{La}_2(\text{CN}_2)_3$ was indexed in the space group I2/a (No. 15) using N-Treor09 ⁶³ and the structure was solved by direct methods (both included in the program EXPO⁶⁴). Other possible candidate space groups suggested by EXPO were lower in symmetry (structure solutions with space groups Ia or I2 were discarded by using the program missym included in the program package PLATON⁶⁵) or gave no reasonable structure solution (I2/m, Im). The crystal structure refinement was carried out with Winplotr (Fullprof), with the final full refinement plot displayed in Fig. 4.

Single-crystal X-ray diffraction

Suitable crystals were selected and mounted on XtaLAB Synergy, Dualflex, HyPix diffractometer. The crystals were kept at a steady T=150.0(2) K during data collection. The structure was solved, and the space group was determined with the ShelXT 2018/2 (Sheldrick, 2018) solution program using dual methods and by using Olex2 1.5-ac5-024 ⁶⁶ as the graphical interface. The model was refined with ShelXL 2018/3 ⁶⁷ using full matrix least squares minimization on F^2 . All non-hydrogen atoms were refined anisotropically. Hydrogen atom positions were found in the electron difference map and refined.

Infrared spectroscopy

The infrared spectroscopy was performed using a Bruker VERTEX 70 Fourier-transform infrared (FT-IR) spectrometer, with the measurement recorded in the spectral range of 400–4000 cm⁻¹. The samples were prepared as potassium bromide (KBr) pellets, with a pure KBr pellet serving as the reference for baseline correction.

Photoluminescence spectroscopy

Emission and excitation spectra of $La_2(CN_2)_3$: Ce^{3+} were measured using an FLS920 fluorescence spectrometer, Edinburgh Instruments, equipped with a 450 W xenon arc lamp OSRAM. The sample chamber was fitted with a mirror optic, specifically designed for powder samples. Detection was carried out with a R2658P single-photon counting photomultiplier tube Hamamatsu. The luminescence spectra were recorded at a spectral resolution of 1 nm, dwell time 0.5 seconds per 1 nm step, and the measurement was repeated three times. The correction file for the emission spectra was obtained from calibration with a tungsten incandescent lamp

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certified by NPL (National Physics Laboratory, UK). For recording temperature dependent emission spectra, a cryostat "MicrostatN" from Oxford Instruments was introduced in the above described spectrometer. The decay curves were measured using an Edinburgh Instruments FLS920 fluospectrometer with a photomultiplier rescence Hamamatsu H74422-40. The samples were excited with an Edinburgh EPL655 pulsed diode laser (65 ps pulse width).

Author contributions

H.-J. M.: conceptualization, supervision, funding acquisition, writing, review, and editing. P. S.: synthesis, PXRD, IR, TGA, writing. E. B.: writing, preparation, experimental analysis. M. S.: X-ray diffraction refinements and structure solutions. T. J. and D. E.: photoluminescence spectroscopy. All authors have read and agreed to the published version of the manuscript.

Data availability

Crystallographic data have been deposited at the CCDC under 2403637 (1), 2409323 (2), and 2393819 ($La_2(CN_2)_3$).†

Data are available within the article.

The data that support the findings of this study are available on request from the corresponding author, H.-J. Meyer.

Conflicts of interest

The authors declare no conflict of interest.

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