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Introduction

White light production is important in a modern society and has applications in ambient lighting and/or screen backlighting in liquid crystalline displays. Presently, light emitting diodes (LEDs) are becoming the state of the art in light production as they can be highly efficient and tunable.

To obtain white light emission (WLE), multiple colours need to be combined on a single device since LEDs are usually monochromatic emitters.¹ Hence, dichroic, trichroic and tetrachroic approaches have been used that combine blue/yellow, red/green/blue (RGB) or RGB/cyano emitters, respectively.² The use of multiple colour components implies a signicant difference in comparison to broadband emitters, such as the

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White light production is of major importance for ambient lighting and technological displays. White light can be obtained by several types of materials and their combinations, but single component emitters remain rare and desirable towards thinner devices that are, therefore, easier to control and that require fewer manufacturing steps. We have designed a series of dysprosium(III)-based luminescent metallacrowns (MCs) to achieve this goal. The synthesized MCs possess three main structural types $\text{LnGa}_4(L')_4(L'')_4$ (type A), $\text{Ln}_2\text{Ga}_8(L')_8(L''')_4$ (type B) and $\text{LnGa}_8(L')_8(OH)_4$ (type C) (H₃L', HL'' and H₂L''' derivatives of salicylhydroxamic, benzoic and isophthalic acids, respectively). The advantage of these MCs is that, within each structural type, the nature of the organic building blocks does not affect the symmetry around Dy³⁺. By detailed studies of the photophysical properties of these Dy³⁺-based MCs, we have demonstrated that CIE coordinates can be tuned from warm to neutral to cold white by (i) defining the symmetry about Dy^{3+} , and (ii) choosing appropriate chromophoric building blocks. These organic building blocks, without altering the coordination geometry around Dy^{3+} , influence the total emission profile through changing the probability of different energy transfer processes including the ${}^{3}T_1 \leftarrow Dy^{3**}$ energy back transfer and/or by generating ligand-centered fluorescence in the blue range. This work opens new perspectives for the creation of white light emitting devices using single component tetrachroic molecular compounds. EDGE ARTICLE
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sun. Generally, the broader the emission profile, and the more colour components, the better is the colour rendition.³

The CIE (Commission Internationale de l'Eclairage) colour space is commonly used to standardize how the average human eye will perceive the colour of an object.⁴ The CIE chromaticity coordinates are defined by three numbers (x, y, z) . The chromaticity of an object can be plotted in a two-dimensional CIE diagram, since it implies a normalization condition: $z = 1 - x \nu$. The white light region lies in the centre of the diagram, with the neutral white light having coordinates (0.33, 0.33). White light can also be described by its Correlated Colour Temperature (CCT) , where its quality is defined by the blackbody equivalent temperature. Modern white light emitting devices include sources such as fluorescent lamps,⁵ high-pressure sodium-mercury lamps,⁶ solid state light emitting diodes,⁷ organic light emitting diodes (OLEDs).^{8,9}

Single component white light emitters are desirable because they allow for thinner materials with greater manufacturing reproducibility and enhanced stability over time.^{10,11} Several lanthanide(III) ions (Ln^{3+}) emitting in the visible range, namely, Sm^{3+} , Eu³⁺, Tb³⁺, Dy³⁺, Tm³⁺, have the potential to emit compatible lights. Typically, the creation of individual materials that display WLE involves various Ln^{3+} ions, each with different properties, either independently or in combination with the

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[†] Electronic supplementary information (ESI) available: Detailed of the synthesis and characterization of MCs, additional information about photophysical studies, details about the determination of the singlet, triplet states, CIE 1931 coordinates and CCT. See DOI: <https://doi.org/10.1039/d4sc00389f>

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light emitted by organic chromophoric ligands or an inorganic matrix.

Several compounds possessing WLE capabilities have been created with a combination of different visible-emitting Ln^{3+} . The variation of the concentrations of the individual $Ln³⁺$ allows one to tune the corresponding relative emission intensities and, thus, to control the CIE coordinates. The pertinence of this strategy has been demonstrated for molecular materials,^{12,13} Ln^{3+} -based polymers,^{14–21} metal–organic frameworks $(MOFs),²²⁻²⁶$ inorganic materials,²⁷⁻²⁹ and hybrid organic/ inorganic³⁰⁻³³ materials.

Alternatively, to produce WLE, the combination of emission signals arising from a single visible-emitting Ln^{3+} and organic ligands or inorganic matrix can be used.34,35 The validity of this approach has been demonstrated for molecular Eu^{3+} compounds,³⁶ as well as for Dy^{3+} compounds. Some of these Dy^{3+} compounds were shown to be tunable by ligand modifications, 37 as well as by excitation wavelength³⁸ or by the change of the experimental temperatures.³⁷ The appearance of the Ln^{3+} emission is dependent on the environment around Ln^{3+} that is controlled by the ligand field induced by the structure of this ligand. It will control the splitting of the different transitions as well as their relative emission intensities and, therefore, the CIE coordinates.³⁹–⁴² Chemical Science

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Single component Dy³⁺ is a special lanthanide(m) ion for such application as its possesses two main groups of sharp emission bands in the green and in the red spectral domains that allow for the development of single-component WLE with the addition of a blue emission component arising from the same compound.⁴⁶⁻⁵⁵ Several inorganic compounds have shown an ability to tune the CIE coordinates by controlling the concentration of the Dy³⁺ dopant. These examples include Dy³⁺doped chloroborosilicate glasses,⁵⁶ barium silicate,⁵⁷ zincaluminum-sodium-phosphate,⁵⁸ or lithium-zinc borosilicate⁵⁹ glasses.

The most studied transitions in Dy^{3+} compounds with respect to WLE⁴⁶ are bands located in the visible range, in particular, ${}^{4}F_{9/2} \rightarrow {}^{6}H_{15/2}$ (blue), ${}^{4}F_{9/2} \rightarrow {}^{6}H_{13/2}$ (yellow), minimally ${}^{4}F_{9/2} \rightarrow {}^{6}H_{11/2}$ (red), and rarely ${}^{4}F_{9/2} \rightarrow {}^{6}H_{9/2}$ (red-brown) transitions. The emission intensity ratio $I(^4\mathrm{F}_{9/2} \rightarrow \mathrm{^6H_{13/2}})/I(^4\mathrm{F}_{9/2}$ \rightarrow $^{6}\mathrm{H}_{15/2})$ is sometimes used as a measurement of the effects of ligand field on the emission spectrum of Dy^{3+} (the "yellow/blue" or Y/B ratio). Modification of the Y/B ratio has been associated with Dy^{3+} -ligand covalency,^{40,41} as well as with asymmetry of the coordination environment.58,60–⁶³ Higher level of covalency and coordination sphere asymmetry apparently leads to a higher Y/B ratio, *i.e.* change of the Dy³⁺ emission profile and tuning of WLE properties.

The coordination environment around $Ln³⁺$ impacts its emission profile and, in turn, CIE coordinates. $Ln³⁺$ -doped extended solids, amorphous inorganic compounds, such as ceramics, glasses, sol-gels or cementitious materials do not possess a single well-defined structure about each dopant.⁷ In crystalline materials, the dopants are dispersed statistically at allowed positions within the lattice, while the introduction of the dopant itself will induce the distortion of the structure of the lattice.⁶⁴–⁶⁷ Thus, in such materials, the correlation between

the emission spectra of the doped Ln^{3+} with the symmetry about them is limited, as changes will be averaged by the environments around each dopant.⁶¹

Single Ln^{3+} molecular compounds provide the advantage of a unique and well-defined coordination environment around the lanthanide, especially when highly rigid. They have been studied for WLE in the past. For example, a single molecular white light emitting Eu^{3+} complex has been previously examined.³⁶ This complex was formed with a 1,8-naphthalimide functionalized tetraazacyclododecane ligand. In this complex, the 1,8-naphthalimide chromophoric ligand generates a blue/ purple fluorescent component, as well as a green component from a naphthalimide aggregation associated excimer state. This ligand also acts as a sensitizer for the generation of the red emission from Eu^{3+} . Through their combination, these different components lead to the generation of WLE.

 Dy^{3+} -based molecular compounds have also shown a strong potential as single-component white light emitters. Thus, by combining Dy^{3+} -centred electronic transitions with those located on the blue-emitting ligand, WLE was observed in a $[Dy^{3+}(TETP)(NO₃)₃]\cdot 4H₂O MOF material [TETP = 1,1',1".$ $((2,4,6\text{-}triethyl\text{-}benzene-1,3,5\text{-}triv)tris(methylene))tris(pyridin-$ 4(1H)-one))]. The CIE coordinates of this MOF could be tuned to $(0.33, 0.35)$ by varying the excitation wavelength.³⁸

In a series of molecular binuclear Dy^{3+} complexes, $[Dy_2]$. $\text{L})_2(\text{NO}_3)_2(\text{solvent})_2] \cdot x(\text{solvent}) \text{ [L = 2,2'-[[(2-pyridinyl-methyl)]$ imino]di(methylene)]bis(4 R-phenol), where $R = CH_3$, Cl, and $CH₃O$; solvent = methanol or DMF], the WLE could be tuned by varying both the substituents of the ligand and the experimental temperature.³⁷ This thermal behaviour might be explained by the interactions of the Dy^{3+} with solvent vibrational modes and/or changes in energy transfer from the ligand. On the other hand, the influence of the substituents of the ligand on the spectra of Dy^{3+} was not fully elucidated but the modification of the CIE coordinates may be attributed to changes of the relative intensities of Dy^{3+} - vs. ligand-centred emissions.

Molecular compounds are more suited probes of the effects of the ligand field on Dy^{3+} emission profile as they provide a defined environment about the coordinated ion. In a recent communication, we have demonstrated that, within a series of monomeric and dimeric Dy^{3+}/Ga^{3+} metallacrowns (MCs) with a common [12-MC_{GaN(Shi)}-4] motif, the Dy³⁺ emission profile can be tuned by modifying the symmetry about the Dy^{3+} ion.⁷⁰ The changes of the relative intensities of different Dy^{3+} emission bands combined with the minimal ligand contribution led to tunable WLE arising from these MCs. Here, we expand this series of MCs with WLE properties by synthesizing eleven new Dy^{3+} MCs formed with different combinations of building blocks using substituted salicylhydroxamic, benzoic and isophthalic acids (Fig. 1) while preserving three main structural types: LnGa₄(L')₄(L'')₄ (type A), Ln₂Ga₈(L')₈(L''')₄ (type B) and LnGa $_8$ [L $')_8$ (OH) $_4$ (type C) (Fig. 1, H $_3$ L $'$ = salicylhydroxamic acid (H₃shi), 5-methylsalicylhydroxamic acid (H₃mshi), 5-methoxysalicylhydroxamic acid (H3moshi), 5-chlorosalicylhydroxamic acid (H₃Cl-shi), 5-bromosalicylhydroxamic acid (H₃Br-shi), 5iodosalicylhydroxamic acid (H₃I-shi); $HL'' = \text{benzoic acid}$

Fig. 1 (A) Schematic presentation of a Dy³⁺[12-MC_{GaN(R}1_{-shi)}-4] MC core, where Ga³⁺ and R¹-substituted H₃shi (B) template the formation of the MC, while Dy³⁺ is linked to the central cavity by the derivatives of benzoic or isophthalic acids (C). (D–F) Top-down (left) and side-on (left-centre) views of the three Ga³⁺/Dy³⁺ MCs geometries and the corresponding primary coordination spheres about Dy³⁺ (centre-left and right views, the red lines between oxygen atoms are guides for the eyes to suggest O₄ planes coming from the MC ring and otherwise). (D) [DyGa₄(shi)₄(bz)₄] representing the LnM₄(L')₄(L'")₄ structure (type A).⁴³ (E) [Dy₂Ga₈(shi)₈(iph)₄] corresponding to the Ln₂M₈(L')₈(L'")₄ structure (type B).⁴⁴ (F) [DyGa₈(shi)₈(OH)₄] corresponding to the LnM₈(L')₈(OH)₄ structure (type C).⁴⁵ The structure was solved for isostructural [NdGa₈(shi)₈(OH)₄]Na. Solvents of crystallization, counter ions, and hydrogen atoms are omitted for clarity. Central Ln³⁺ and coordinating oxygen atoms are bolded for highlight effect. Colour code: Ga, orange; Ln, teal; O, red; N, blue; C, grey.

(Hbz), 2-methylbenzoic acid (Hmbz), 2-fluorobenzoic acid (HFbz); $H_2L''' =$ isophthalic acid (H₂iph), 5-iodoisophthalic acid (H2I-iph)). This approach allows the investigation of the influence of these modifications to the electronic structure of the ligand scaffold may have on the total emission profile of Dy^{3+} MCs and on the corresponding WLE properties reflected by the CIE coordinates. Moreover, to determine how such modifications affect the energy positions of the singlet and triplet states within this series of MCs, the corresponding Gd^{3+} analogues were synthesized and studied.

Experimental section

 Ln^{3+}/Ga^{3+} MCs were synthesized using the deprotonated forms of the ligands given in Fig. 1. All MC synthesis reactions were carried out aerobically under ambient conditions via single-pot self-assembly reactions. Elemental analyses were performed by Atlantic Microlabs Inc. ESI-MS spectra were collected with an Agilent 6230 TOF HPLC-MS mass spectrometer in negative ion mode (−350 V) on samples dissolved in methanol at a concentration of 2 $\mathrm{mg}\:\text{mL}^{-1}\text{.}^1$ H NMR spectra were collected on a Varian MR400 NMR in deuterated DMSO at a concentration of 4 mg mL⁻¹. Single crystal unit cell parameters were obtained by mounting samples on a Rigaku AFC10K Saturn 944+ CCD-based X-ray diffractometer equipped with a low temperature device and Micromax-007HF Cu-target micro-focus rotating anode ($\lambda =$ 1.54187 Å) operating at 1.2 kW power (40 kV, 30 mA). The X-ray intensities were measured at 85(1) K with the detector placed at a distance of 42.00 mm from the crystal. Detailed synthetic procedures and characterizations are provided in ESI.† Photophysical characterization was carried out as described in ESI.†

Results

Synthesis and characterization

 Ln^{3+}/M^{3+} MCs possess the remarkable property to form structurally closely similar compounds when the ligands used for the assembly possess identical binding motifs. Furthermore, by modifying the binding motif, the coordination geometry can also be tightly controlled and modified. Thus, independently of the nature of the $Ln^{3+} (Ln^{3+} = Pr^{3+} - Yb^{3+})$ and for a variety of M^{3+} metals located in the ring $(M^{3+} = Al^{3+}, Ga^{3+}, Mn^{3+}, Fe^{3+})$, the use of the salicylhydroximate 'O–C–C–CO–NHOH' motif in combination with carboxylic or hydroxy ligands has been shown to form similar monomeric $\text{LnM}_4(L')_4(L'')_4$ (Fig. 1D), $^{43,73-76}$ dimeric $\rm Ln_2M_8(L')_8(L''')_4$ (Fig. 1E)^{44,71,77–79} or $\rm LnM_8(L')_8(OH)_4$ (Fig. 1F)⁴⁵ structural topologies, respectively.

Novel functionalized salicylhydroxamic acids were synthesized in this work by converting the appropriate carboxylic acids into the corresponding methyl esters followed by an exchange reaction with an excess of hydroxylamine. The MCs were synthesized by a self-assembly reaction between stoichiometric amounts of derivatives of salicylhydroxamic, benzoic or isophthalic acids with Ga^{3+} and Ln^{3+} nitrates in DMF or methanol in the presence of a base. MCs were obtained as pure crystalline products by the slow evaporation of the solvent. The composition of the MCs was confirmed by mass spectrometry and elemental analysis (ESI†). Each novel MC was screened for unit cell parameters by single crystal X-ray diffraction to confirm the crystallinity of the sample and to establish the crystallographic similarities for different Dy^{3+} and Gd^{3+} analogues. Several novel Dy^{3+} and Gd³⁺ MCs were synthesized using this approach. For the sake of comparison, the spectroscopic properties of these

compounds are analysed and presented together with those obtained from several Dy^{3+} and Gd^{3+} MCs which we previously described.⁴³–45,68,69

In total, eight of these complexes adopt the $\mathrm{LnGa}_4(\mathrm{L}')_4(\mathrm{L}'')_4$ topology (Type A), five adopt the $\text{Ln}_2\text{Ga}_8(\text{L}')_8(\text{L}''')_4$ topology (Type B), and two adopt the $\text{LnGa}_8(\text{L}')_8(\text{OH})_4$ topology (Type C). Most

Fig. 2 Corrected and normalized emission spectra of the Dy^{3+} MCs in the solid state upon ligand-centred excitation at 340 nm at room temperature. Counter-cation is either Na⁺ or 2 Na⁺ unless stated otherwise.

Table 1 Photophysical data for the Dv^{3+} MCs under ligand-centred excitations at room temperature^a

of these MCs crystallize with $Na⁺$ as counterions, while two possess a PyH $^{\scriptscriptstyle +}$ (pyridinium) counterion and one include NH $_4^{\scriptscriptstyle +}$ counterions. Throughout this article, the counter-cations are $Na⁺$ or 2 Na⁺ unless stated otherwise. Each compound herein presented also co-crystallizes with a variable number of solvent molecules (DMF or methanol, and water). The details about the synthesis and the characterization of all studied MCs are given in the ESI.†

Photophysical properties

The photophysical properties of Dy^{3+} MCs were measured in the solid state at room temperature. Each of the complexes showed a characteristic Dy^{3+} emission profile under ligand excitation at 340 nm (Fig. 2). These profiles include three emission bands, two, being the most intense, correspond to the ${}^{4}F_{9/2} \rightarrow {}^{6}H_{15/2}$ (\sim 480 nm, blue) and ⁴F_{9/2} → ⁶H_{13/2} (\sim 575 nm, yellow) transitions, and one, being significantly less intense, is due to the $^4\mathrm{F}_9$ ₂ → ⁶H_{11/2} (~660 nm, red) transition. A broad band located in the blue is attributed to the emission arising from the electronic structure of the chromophoric building blocks that constitute the different MCs. Important differences in intensities of these blue bands in comparison to the intensities of the Dy^{3+} transitions can be observed in the emission spectra (Fig. 2).

Absolute quantum yields of the Dy³⁺-centred (Q_{Dy}^L) and ligand-centred (Q_{L}^L) transitions were determined upon excitation in the range 330–350 nm. The total quantum yield value (Q_{total}^L) was calculated using the formula: $Q_{\text{total}}^L = Q_{\text{Dy}}^L + Q_{\text{L}}^L$. The values of Q_{Dy}^L vary from 0.128(4)% to 8.3(3)% (Table 1), while those of Q_{L}^{L} change from 0.0043(1)% to 0.607(3)%. The experimental decay of the luminescence of monomeric $DyGa_4(L')_4(L''$ ₄ are best described by a monoexponential function while those of $\text{LnGa}_8(\text{L}')_8(\text{OH})_4$ and most of $\text{Ln}_2\text{Ga}_8(\text{L}')_8(\text{L}''')_4$ exhibits a biexponential behaviour. The observed luminescence

^a Data for powder solid state samples of MCs. 2 σ values are given between parenthesis. Estimated experimental errors: $\tau_{\rm obs}$, $\pm 2\%$; $Q_{\rm Dy}^L$, $\pm 10\%$; $\tau_{\rm obs}$ +10%; $\sigma_{\rm sys}^L$, $\pm 10\%$; $\tau_{\rm obs}$ and population parameters $P_i = \frac{B_i \tau_i}{\sum_{i=1}^{n} B_i}$ $\sum_{i=1} B_i \tau_i$ in % are given after the colon. d Under excitation at 340 nm. e Quantum yield of Dy³⁺-centred transitions.

 f Quantum yield of ligand-centred transitions. g $Q^L_{\text{total}} = Q^L_{\text{Dy}} + Q^L_{\text{Lz}}$

^a Data for powder solid state samples of MCs. ${}^{3}T_1$, S₁, and ΔE values are given in cm⁻¹. The energy values of ${}^{3}T_1$ and S₁ determined using an alternative method are provided in Table S2. b Counter-cation is Na⁺ or 2 Na⁺ unless otherwise specified. ^c Determined from the Gaussian fitting of the 0–0 phonon line of the phosphorescence spectra of Gd^{3+} MCs measured at 77 K (Fig. S6, ESI). d Determined from the red edge (10% of the maximum) of the solid-state diffuse reflectance spectra (Fig. S2, ESI). e Energy of the emissive ${}^{4}F_{9/2}$ level of Dy³⁺ is 21 050 cm⁻¹⁷. \hat{f} Gd³⁺ analogue could not be synthesized, so ${}^{3}T_1$ is not determined. g Determined for Dy³⁺ analogues.

lifetimes of the Dy³⁺ emission vary from 4.1(1) to 61.1(7) µs. The excitation spectra are given in Fig. S1 (ESI).†

The diffuse reflectance spectra of Dy^{3+} and Gd^{3+} MCs were measured in the solid state at room temperature (Fig. S2, ESI†). The lowest excited singlet state (S_1) energy levels were estimated by considering the red edge of these spectra (Table 2) or from the intersection of the emission spectrum of the ligand with the diffuse reflectance spectrum of the corresponding MC (Fig. S4, ESI†). The shapes of these two types of spectra match well with one another (Fig. S8, ESI†), and are summarized in Table 2. Energy values vary from $25\,870$ to $28\,860$ cm^{-1} . The corresponding measurements were also taken for the organic building blocks used to assemble MCs (Fig. S3, S5 and Table S2, ESI†).

The phosphorescence spectra were collected at 77 K for all the synthesized Gd^{3+} MCs in the solid state. The blue side of the observed phosphorescence spectrum was deconvoluted into several Gaussian curves. The highest-energy band was taken as the 0–0 phonon phosphorescence emission and considered as the energy of the lowest ligand triplet state $({}^{3}T_{1})$. These values vary from 21 570 to 22 680 $\rm cm^{-1}$. Another estimate of the $^3\rm{T}_1$ energy level was also calculated as the centre of gravity of the phosphorescence spectrum. These values are presented in Table S2 (ESI†) and have a similar trend as the ones obtained from the 0–0 triplet state energy estimation (Fig. S8, ESI†). When sufficient emission signals could be obtained, the phosphorescence spectra of the free ligands were acquired in the solid state and analysed (Fig. S7 and Table S2, ESI†). The ligands singlet and triplet states are presented relatively to the relevant states of Dy^{3+} in Fig. 3.

Colorimetric properties. The colorimetric properties of the Dy^{3+} MCs were calculated from the corresponding visible emission spectra in the range of 370–700 nm (Fig. 2 and ESI†).⁸⁰ Each MC showed CIE coordinates consistent with WLE, with x

Fig. 3 Diagram of Dy^{3+} energy levels (black traces), the lowest singlet $(S_1,$ dotted traces) and triplet $({}^3T_1,$ solid traces) states determined for Gd³⁺ MCs as well as for the organic building blocks used to assemble MCs. The list of energy levels is provided in Tables 2 and S2, ESI.† Chemical structures of the ligands and crystal structures of MCs are given in Fig. 1.

coordinates ranging from 0.29 to 0.35 and y coordinates ranging from 0.28 to 0.39 as compared to the neutral WLE coordinates of x, $y = (0.33, 0.33)$ (Table 3). These results demonstrate the possibility to tune the WLE of Dy^{3+} MCs within the studied series based principally on the nature of the ligands and, to a lower extent, to the symmetry around the Dy^{3+} (Fig. 4). The CIE diagram is presented separately for each MC structure type in Fig. S11 (ESI).†

For the type A structures, each of the Dy^{3+} MCs possesses similar coordinates (0.34–0.35, 0.38–0.39). However, two of the compounds, $DyGa_4(Cl-shi)_4(bz)_4$ and $DyGa_4(Cl-shi)_4(F-bz)_4$ have disparate CIE coordinates (0.32, 0.33). Similarly, in the case of the type B structures, each Dy^{3+} MC has similar CIE coordinates $(0.33-0.35, 0.36-0.39)$ except for the $Dy_2Ga_8(moshi)_8(iph)_4$ which has disparate coordinates (0.29, 0.28). For the type C structures, each of the two compounds has similar coordinates (0.31–0.32, 0.33–0.34).

Emission spectra of Dy^{3+} MCs, beside the emission signals due to the ${}^{4}F_{9/2} \rightarrow {}^{6}H_J (J = 15/2, 13/2, 11/2)$ Dy³⁺ transitions, exhibit broad ligand-centred emission bands possessing different intensities in the UV-blue range (Fig. 2). To estimate if ligandcentred emission signals affect CIE coordinates, they were recalculated for the wavelength range comprised between 460 and 700 nm that covers exclusively the Dy^{3+} -centred transitions (Table 3). The values obtained are much more similar for each MC structure type, i.e. (0.35–0.36, 0.39–0.40) for type A, (0.34–0.36, 0.39–0.40) for type B, and (0.32–0.33, 0.36–0.37) for type C, suggesting that the Dy^{3+} emission is influenced more by the symmetry of the coordination sphere rather than by the nature of the coordinating ligand. This is further confirmed by the small variations of the relative integral intensities of different Dy^{3+} transitions observed in the emission spectra, in particular the ratio of the yellow (${}^{4}F_{9/2} \rightarrow {}^{6}H_{13/2}$) to the blue (${}^{4}F_{9/2} \rightarrow {}^{6}H_{15/2}$) transitions, within each MC structural type (Tables 3 and S1†).

Fig. 4 CIE 1931 diagram depicting the chromaticity coordinates (calculated for the 370–700 nm range) for all of the Dy³⁺ MCs presented herein. The CIE coordinates are specified in Table 3 and presented for each structure type separately in Fig. S11, ESI.†

Discussion

Fifteen Dy^{3+} MCs formed with varied organic building blocks that belong to three different molecular topologies (Fig. 1) were synthesized and characterized. Their photophysical properties in the solid state, *i.e.* diffuse reflectance, excitation and emission spectra, observed luminescence lifetimes, ligand- and Dy^{3+} -centred quantum yields were acquired and analysed. CIE 1931 and CCT have been calculated. Several Gd^{3+} analogues were also synthesized to provide complementary information about the ligand-centred electronic properties. This large array of compounds allows a fairly comprehensive exploration of whether the introduction of different substituents to the MC constituting ligands in a series of Dy^{3+}/Ga^{3+} MCs adopting three different molecular topologies can influence the emission profiles toward the creation of tunable molecular-based WLE diodes.

Among different visible-emitting Ln^{3+} -based materials, Dy^{3+} ones are unique in their ability to produce WLE.⁴⁶⁻⁵⁵ Indeed, a typical emission spectrum of Dy^{3+} in the visible range contains three fundamental components: a blue, a yellow and a red one arising from the ${}^{4}F_{9/2}$ → ${}^{6}H_{15/2}$ (~480 nm), ${}^{4}F_{9/2}$ → ${}^{6}H_{13/2}$ (~575 nm), and ${}^{4}F_{9/2} \rightarrow {}^{6}H_{11/2}$ (~660 nm) electronic transitions, respectively. Therefore, the WLE of the material can be tuned by modifying the Dy^{3+} emission profile and the relative intensities of each of these transitions. These parameters, in turn, can be controlled by the nature of the surrounding environment (in this case organic ligands) and the coordination symmetry about Dy^{3+} . In addition, organic ligands may exhibit emission in the visible range and contribute to the CIE coordinates. Indeed, the generally accepted sensitization mechanism of $Ln³⁺$ in coordination compounds is assumed to be occurring through an energy transfer from the ${}^{3}\mathrm{T}_1$ (mainly) and/or S_1 states of organic ligands, i.e. 'antenna effect'. If this energy transfer is not complete, emission spectra may exhibit residual broad bands arising from the electronic structure of the ligand along with Ln^{3+} -centred transitions. Therefore, the analysis of the effect of the nature of organic ligands on the emission spectra of Dy^{3+} based compounds is key to rationalize and further tune the WLE properties.

If one considers the ligand-centred photophysical properties in the studied series of MCs, more pronounced changes of the diffuse reflectance spectra (Fig. S3, ESI†), emission profiles (Fig. S4, ESI†) and energy positions of ${}^{3}T_{1}$ and/or S₁ states (Table 2 and Fig. 3) are observed upon introduction of different substituents to the salicylhydroximate ligands, H_3L' . For example, ${}^{3}T_{1}$ and S_{1} are lowered by 1060 and 915 $\rm cm^{-1}$ for $\left[\text{GdGa}_{4}(\text{Cl-shi})_{4}(\text{bz})_{4}\right]$ or by 745 and 2780 cm⁻¹ for $\left[\text{Gd}_{2}\text{Ga}_{8}(\text{Hilj})_{4}(\text{dz})_{4}\right]$ moshi)₈(iph)₄] compared to $\left[GdGa_4(\text{shi})_4(\text{bz})_4 \right]$ or $\left[Gd_2Ga_8(\text{-pb})_4 \right]$ shi)8(iph)4], respectively. Accordingly, the maxima of the ligandcentred emission bands are red-shifted by 17 nm (from 360 nm for $\left[\text{GdGa}_{4}(\text{shi})_{4}(\text{bz})_{4}\right]$ to 377 nm for $\left[\text{GdGa}_{4}(\text{Cl-shi})_{4}(\text{bz})_{4}\right]$ or 39 nm (from 364 nm for $\left[\frac{Gd_2Ga_8(\text{shi})_8(\text{iph})_4}{\text{pth}} \right]$ to 403 nm for $[\text{Gd}_2\text{Ga}_8(\text{moshi})_8(\text{iph})_4]$. If one compares the diffuse reflectance spectra of the LnGa₄(L')₄(L'')₄, Ln₂Ga₈(L')₈(L''')₄ and LnGa₈(- $\rm L^{\prime})_{8}(OH)_4$ MCs $\rm (Ln ~ = ~Gd^{3+} ~or ~ Dy^{3+})$ with those of the

corresponding constituent organic building blocks (Fig. S9, ESI†), the positions of the red edges of the spectra and, in turn, the S_1 levels (Fig. 3 and Table S2, ESI[†]), are mainly determined by the nature of the ring salicylhydroximate ligands, $H₃L'$. On the other hand, a similar comparison of the ligand-centred emission profiles (Fig. S10, ESI†) is more challenging. Generally, the maxima of the emission bands of the Gd^{3+} MCs are redshifted by 20–40 nm compared to those of the ring H_3L' ligands. However, in the case of $\left[GdGa_4(Br\text{-}shi)_4(bz)_4 \right]$ and $\left[Gd_2Ga_8(-f)\right]$ $moshi|_8(iph)_4$, a blue shift of the emission maxima by 40 or 52 nm is observed, respectively, to these of $H_3Br\text{-}shi$ or H₂moshi.

The emission spectra of all Dy^{3+} MCs exhibit three main bands in the visible range originating from the ${}^4\mathrm{F}_{9/2} \rightarrow {}^6\mathrm{H}_J$ (*J* = 15/2, 13/2, 11/2) electronic transitions along with the broad ligand-centred emission < 460 nm (Fig. 2 and Table 1). The relative intensity of the latter depends on the nature of the constituting organic ligands and is the most pronounced for $[DyGa_4(Cl-shi)_4(bz)_4]$, $[DyGa_4(Cl-shi)_4(F-bz)_4]$ and $[Dy_2Ga_8({\rm mod}(i\pi)$ compared to ${\rm [DyGa_4(shi)_4(bz)_4]}$ or ${\rm [Dy_2Ga_8]}$. $\frac{\sinh\sinh(1)}{\sinh(1)}$. On the other hand, relative integral intensities (Table S1, ESI†) and the ligand field splitting of the different Dy^{3+} transitions are determined by the symmetry about Dy^{3+} . Moreover, relative integral intensities are similar for Dy^{3+} MCs of type A and B being in line with the analogous coordination environments around Dy^{3+} in both structures (Fig. 1). In contrast, significant differences are observed when comparing the emission spectra of types A and B versus type C, which has a more distinct Dy^{3+} -coordination environment. All these variations, in turn, impact both the WLE properties and CIE coordinates (vide infra). Edge Article

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When it comes to quantitative characteristics, *i.e.* Dy³⁺-centred quantum yield Q_{Dy}^L and observed luminescence lifetimes (Table 1), more pronounced changes are again observed upon introduction of different substituents to the salicylhydroximate ligands. More specifically, for the halogenated analogues of Dy³⁺ MCs of type A the values of $\tau_{\rm obs}$ are shorter by 4.5–5.9 times, while Q_{Dy}^{L} are lower by 9.7–21 times compared to those of the $[DyGa_4(\text{shi})_4(\text{bz})_4]$ MC. If one considers Dy^{3+} MCs of type B, the introduction of a $-OCH₃$ substituent on the salicylhydroximate ligand leads to a 7.4-times shortening of the longest component of $\tau_{\rm obs}$ and 11.4-times lowering of the $Q_{\rm Dy}^L$ relative to the [Dy₂- $Ga_8(\text{shi})_8(\text{iph})_4$. On the other hand, an introduction of iodine on the isophthalate ligand in $[Dy_2Ga_8(shi)_8(I-iph)_4]$ results in a 2.2-times lengthening of the longest component of $\tau_{\rm obs}$ (up to 40.7 μ s) and an insignificant variation of the value of Q_{Dy}^L . similar value of τ_{obs} (37.7 µs) is observed for [DyGa₈(shi)₈(OH)₄]. These observations correlate well in general with the positions of the ${}^{3}T_{1}$ relative to the ${}^{4}F_{9/2}$ level of Dy³⁺: the smaller the difference between these two levels, the lower is the value of the Q_{Dy}^{L} because of the higher probability of back ${}^{3}T_{1} \leftarrow Dy^{3*}$ energy transfer. We can also mention that the longest values of $\tau_{\rm obs}$ are observed for MCs with a ${}^{3}T_{1}$ state located at least 1000 cm⁻¹ above the Dy³⁺⁴F_{9/2} accepting level and the S₁ state not being in resonance with any upper lying Dy^{3+} states. However, among the Dy^{3+} MCs possessing very similar energy positions of the ${}^{3}T_{1}$, e.g. $[DyGa_4(Cl\text{-}shi)_4(bz)_4]$,

 $[Dy_2Ga_8(moshi)_8(iph)_4]$ and $[Dy_2Ga_8(mshi)_8(iph)_4]$ (Table 2), the values of Q_{Dy}^L vary from 0.128 to 0.837% with the lowest one being for $[Dy_2Ga_8(moshi)_8(iph)_4]$. This observation can be explained by the lower energy position of the S_1 state in this MC and its participation to energy transfer processes, including enhanced back transfer $S_1 \leftarrow {}^{3}T_1 \leftarrow Dy^{3'*}.$ We observed a similar situation for the analogues Th^{3+} MCs.⁶⁹ Enhanced back energy transfer processes also contribute to the relative enhancement of the ligand-centred emission and contribute to the CIE coordinates (vide infra).

WLE properties, CIE coordinates and CCT, of a material are usually determined in the visible range from 370 to 700 nm, where the human eye is sensitive. In the case of the Dy^{3+} MCs studied here, this range includes the broad ligand-centred bands (360–460 nm) and the three Dy³⁺ transitions, ${}^{4}F_{9/2} \rightarrow {}^{6}F_{1}$ and ${}^{4}F_{1} \rightarrow {}^{6}F_{1}$ (460–700 nm) CIE $H_{15/2}$, ${}^{4}F_{9/2} \rightarrow {}^{6}H_{13/2}$ and ${}^{4}F_{9/2} \rightarrow {}^{6}H_{11/2}$ (460–700 nm). CIE coordinates (x, y) lie in the range $(0.32-0.35, 0.33-0.39), (0.29-0.35, 0.33)$ 0.35, 0.28–0.39) and $(0.31-0.32, 0.33-0.34)$ for Dy³⁺ MCs of type A, B and C, respectively (Table 3). Each of these coordinates correspond to a type of white appearing light, ranging from blue (or cool) to neutral (0.33, 0.33) and yellow (or warm) white (Fig. 4). This range represents an attractive tunability of the WLE in these series of Dy^{3+} MCs, controlled by the nature of the constituting organic ligands. The closest to the neutral white light CIE coordinates are observed for $[DyGa_4(Cl-shi)_4(bz)_4]$, $[DyGa_4(Cl-shi)_4(F-bz)_4]$ and Dy^{3+} MCs of type C.

Since the emission spectra of Dy^{3+} MCs originate from both the Dy³⁺ and the constituting ligands, another set of (x, y) was calculated considering only Dy^{3+} -centred transitions in the range of 460–700 nm (Table 3) in order to understand which parameters and to what extent they contribute to the tunability of CIE coordinates. The obtained values indicate a significantly lower level of tunability: similar for MCs of types A and B while distinct for those of type C, and vary only within 0.02 units for both x and y for a specific MC topology. These results are in line with the previous point that the Dy^{3+} emission profile, *i.e.* the relative integral intensities of the ${}^4\mathrm{F}_{9/2} \rightarrow {}^6\mathrm{H}_{\mathrm{J}}(\mathrm{J} = 15/2, 13/2, 11/2)$ 2) transitions and their ligand field splitting, is determined by the MC topology and the symmetry about Dy^{3+} , and is less influenced by the nature of the constituent organic ligands. On the other hand, the latter has a signicant impact on the general emission profile and contributes significantly to the tuning of the WLE properties within a series of Dy^{3+} MCs of a specific structure type.

Here, we have demonstrated that, among MCs of types A and B, the greatest divergence from average CIE coordinates, ca. (0.35, 0.39), is observed for $[DyGa_4(Cl-shi)_4(bz)_4]$ (0.32, 0.33), $[DyGa_4(Cl-shi)_4(F-bz)_4]$ (0.32, 0.33), $[Dy_2Ga_8(mshi)_8(iph)_4]$ (0.33, 0.36), and $[{\rm Dy}_2\text{Ga}_8(\text{moshi})_8(\text{iph})_4]$ (0.29, 0.28). The origin of the disparities using these building blocks must come from the higher contribution of the ligand-centred bands to the total emission profile because of the similarity of the Dy^{3+} -centred transitions for these types of MCs. Indeed, the ratio $Q_{\text{L}}^{L}/Q_{\text{total}}^{L}$ is the largest for the selected Dy^{3+} MCs and it increases in the row $[Dy_2Ga_8(mshi)_8(iph)_4]$ (0.237) << $[Dy_2Ga_8(moshi)_8(iph)_4]$ (0.435) \langle [DyGa₄(Cl-shi)₄(F-bz)₄] (0.467) \langle [DyGa₄(Cl-shi)₄(bz)₄] (0.513) (Table 1). In general, these results correlate well with the

positions of the ${}^{3}T_{1}$ states of the ligands relatively to the ${}^{4}F_{9/2}$ level of Dy^{3+} and higher probability of the back ${}^{3}T_{1} \leftarrow Dy^{3+}$ energy transfer that has been discussed above. However, one may notice that the most significant impact on CIE coordinates is observed for the $[Dy_2Ga_8(moshi)_8(iph)_4]$ MC, that does not possess the highest $Q_{\text{L}}^L/Q_{\text{total}}^L$ ratio. On the other hand, $[\text{Dy}_2$ - Ga_8 (moshi)₈(iph)₄] exhibits the most red-shifted ligand-centred emission among the studied MCs (Fig. 2), that is located in the blue range of the spectrum and, in turn, has a larger contribution to the CIE coordinates. Indeed, a higher energy (ultraviolet) fluorescence is less visible to the human eye and has an insignificant impact on the tuning of WLE properties of the materials (Fig. S12, ESI†). Therefore, the CIE coordinates of Dy^{3+} MCs of a particular topology can be adjusted by the appropriate choice of the building blocks that will influence the total emission profile by changing the probability of different energy transfer processes including the ${}^{3}T_{1} \leftarrow Dy^{3+*}$ process and/or by exhibiting fluorescence in the blue range. In this way, the warm white emission of the majority of Dy^{3+} MCs of types A and B can be turned into neutral white in $[DyGa_4(Cl-shi)_4(bz)_4]$ (0.32,0.33), and $[DyGa_4(Cl-shi)_4(F-bz)_4]$ (0.32, 0.33), for example. To an extent, this can be predicted before the MC is synthesized, because the S_1 energy of the ligand should generally inversely correlate with the wavelength of the emission, *i.e.* ligands with lower S_1 will exhibit emission at longer wavelengths (Fig. S5, ESI†). However, as mentioned before, the direct correlation of the ligand-centred emission in the MC with the one of the constituent organic ligands is not straightforward. Several different parameters have to be considered. Moreover, in the solid state, additional parameters that will impact the ligandcentred emission need to be taken into account: these include the effect of the presence of the co-crystallized solvent molecules, the intramolecular interactions and interactions due to the packing of the molecules. One should also consider the ratio of the ligand fluorescence to phosphorescence, as a higher proportion of the latter will lead to a more redshifted total ligand-centred emission. Chemical Science

(D₂-Ga_n(modi)₄[iph₁] (Table 2), the measures of the branch of the ²T₁ states of the branch on 1/4/2024 article of C₁-C₂ May 2022. The states of the particle is licensed under the proposit

Conclusions

We have previously shown that Dy^{3+}/Ga^{3+} metallacrowns possessing three different molecular topologies were capable of tuning CIE coordinates through the modification of the Dy^{3+} first coordination sphere symmetry. In this work, we subject these previously characterized systems to electronic perturbations via ligand functionalization to decipher the role of the different organic building blocks on the emission properties of these molecules.

First, we observed that the Dy^{3+} -centred transitions are essentially unperturbed by altering the electron donor or acceptor properties of the organic building blocks. This insensitivity of the Dy^{3+} emission profile to electronic perturbations is a consequence of the rigidity of this family of MCs meaning that one may modify the donor/acceptor properties of the organic ligand of the molecule while retaining the original lanthanide(III) spectral profile. Second, we have demonstrated that CIE coordinates can be extensively tuned from warm to neutral to

cool white by first defining the symmetry about Dy^{3+} and then choosing appropriate organic ligands that will influence the total emission profile through changing the probability of different energy transfer processes including ${}^{3}T_{1} \leftarrow Dy^{3+*}$ energy back transfer and/or by generating ligand-centred fluorescence in the blue range. In particular, the overall WLE profile of the Dy^{3+}/Ga^{3+} MCs can be changed by altering the relative proportion of blue ligand emissions. This behaviour is due to the more visible appearance of the lower energy ligand states' fluorescence (opposed to less-visible higher energy UV emissions), and also the effect that electronic perturbations on the ligands has on the proportion of Dy^{3+} - vs. ligand-centred transitions, $(Q_{\text{L}}^L/Q_{\text{total}}^L)$. Thus, by specifying an individual coordination geometry for the lanthanide(III) in the MC structural motif, one obtains an essentially invariant Dy^{3+} emission that can be fine-tuned to the desired CIE coordinates by controlling the electronic properties of organic building blocks. We have shown that this approach allows the creation of unique tunable WLE using single component tetrachroic, molecular chromophores. Edge Article

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Data availability

All data are available in the ESI.†

Author contributions

E. V. S. – investigation, data curation, formal analysis, writing – original draft; S. V. E. – conceptualization, investigation, data curation, formal analysis, funding acquisition, writing – review & editing; S. P. – conceptualization, funding acquisition, supervision, writing – review & editing; V. L. P. – conceptualization, funding acquisition, supervision, writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

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