Review

Lanthanide Photoluminescence in Heterometallic Polycyanidometallate-Based Coordination Networks

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Received: 16 October 2017; Accepted: 2 November 2017; Published: 4 November 2017

Abstract: Solid-state functional luminescent materials arouse an enormous scientific interest due to their diverse applications in lighting, display devices, photonics, optical communication, low energy scintillation, optical storage, light conversion, or photovoltaics. Among all types of solid luminophors, the emissive coordination polymers, especially those based on luminescent trivalent lanthanide ions, exhibit a particularly large scope of light-emitting functionalities, fruitfully investigated in the aspects of chemical sensing, display devices, and bioimaging. Here, we present the complete overview of one of the promising families of photoluminescent coordination compounds, that are heterometallic d–f cyanido-bridged networks composed of lanthanide(3+) ions connected through cyanide bridges with polycyanidometallates of d-block metal ions. We are showing that the combination of cationic lanthanide complexes of selected inorganic and organic ligands with anionic homoligand \([M(CN)_x]^n^-\) (\(x = 2, 4, 6\) and 8) or heteroligand \([M(L)(CN)_4]^2-\) (\(L = \) bidentate organic ligand, \(M = \) transition metal ions) anions is the efficient route towards the emissive coordination networks revealing important optical properties, including 4f-metal-centred visible and near-infrared emission sensitized through metal-to-metal and/or ligand-to-metal energy transfer processes, and multi-coloured photoluminescence switchable by external stimuli such as excitation wavelength, temperature, or pressure.

Keywords: functional materials; crystal engineering; coordination networks; poly-cyanidometallates; lanthanides; photoluminescence

1. Introduction

Luminescent materials, that are able to emit the light due to absorption of photons, electric current, chemical reactions, or a mechanical action, are applied in the numerous aspects of science, technology, and everyday life. They are utilized in cathode ray or fluorescent tubes, X-ray detectors, lighting and display devices, optical communication, low-energy scintillation, optical storage, light conversion, photovoltaics, chemical sensing, bioimaging, and molecular thermometry [1–6]. Considering the construction of light-emitting devices, there are several particularly desired optical functionalities, including white-light emission (WLE), multi-coloured tunable emission, long-lived near-infrared (NIR) phosphorescence, and the non-linear optical property of up-conversion luminescence (UCL) [7–10]. These functionalities are efficiently realized by the photo- and electroluminescent materials, which are the most attractive in the application aspects [1–6].

Among the functional photoluminescent materials, the prime role has been played by traditional inorganic solids, such as oxides, fluorites, and silicates, that can incorporate emissive metal centres, mainly trivalent lanthanide ions revealing a wide range of photoluminescence ranging from UV for \(\text{Gd}^{3+}\), through visible for \(\text{Eu}^{3+}\) or \(\text{Tb}^{3+}\), to NIR light for \(\text{Nd}^{3+}\) or \(\text{Yb}^{3+}\) [2]. Photoluminescence was also broadly investigated for organic molecules, especially those built of the expanded system of...
aromatic rings [1]. Organic chromophores were applied for the preparation of luminescent lanthanide complexes where the organic part serves as an antenna for sensitization of 4f-centred emission through the energy transfer process [11]. Lately, the increasing interest is devoted to the luminescent metal-organic frameworks (MOFs), that belong to the broader group of luminescent coordination polymers (CPs) [12–14]. MOFs are composed of metal ions or clusters bonded by organic linkers into the three-dimensional coordination networks, while CPs are generally built of metal centres and organic ligands combined infinitely in at least one direction. Metal-organic frameworks have been intensively investigated in the context of their highly porous character of a potential application in hydrogen storage, selective gas adsorption, and gas sensing [15,16]. More importantly, MOFs can reveal diverse ligand-based, metal-based, charge transfer, or guest-induced photoluminescence due to their multi-component structure [13].

As opposed to classical inorganic solid luminophors, the synthesis of luminescent MOFs can be straightforward, often without applying high temperature or pressure. Their structures are predictable, and can be rationally designed through a molecular building block approach. They are transformable into the nanoscale, and can be post-synthetically modified by thermal or chemical treatment. Luminescent MOFs were also found to be efficient platforms for multifunctionality by fruitful addition of magnetic, optical or catalytic functionalities [16]. All these features made MOFs attractive as the luminescent objects, particularly appealing in the aspects of chemical sensing, biomedicine and light-emitting devices [12–14]. Of a special interest are lanthanide-based MOFs which were found to exhibit a large scope of luminescent effects, including white-light and tunable visible emission, as well as NIR and up-conversion luminescence [17–20].

All the positive characteristics of luminescent MOFs can be similarly ascribed to the specific group of coordination systems based on polycyanidometallates. They are constructed of the anionic polycyanidometallate complexes of transition metal ions which are combined through cyanide (CN\(^-\)) molecular bridges with the cationic complexes of various 3d/4d/4f metal ions. Depending on the synthetic conditions, and the addition of supporting organic ligands, they are able to form three-dimensional coordination networks, similar to those characteristic of typical MOFs, but also they can give two- or one-dimensional coordination polymers, and discrete polynuclear clusters of diverse topologies [21–24]. Cyanide bridges offer the close and strong connection between two different neighbouring metal centres which was fruitfully explored in the induction of strong exchange magnetic coupling [25,26]. This feature is attractive for the photoluminescent properties as the short inorganic CN\(^-\) linker can strongly couple two metal centres inducing the efficient metal-to-metal energy transfer process. This sounds promising considering the reported polycyanidometallates of 3d/4d/5d metal ions which were found to be strong absorbers of UV light, exhibiting often weak absorption in the visible and NIR ranges. Such spectroscopic behaviour makes them the great candidates for the sensitization of luminescent trivalent lanthanide ions. In addition, some of the polycyanidometallates of transition metal ions reveal the intrinsic photoluminescent properties related to the metal-centred (MC) d-d transitions, the metal-to-ligand charge transfer (MLCT), or the opposite ligand-to-metal charge transfer (LMCT) effects [27–33].

Heterometallic polycyanidometallate-based coordination networks have been extensively studied in the last decades due to their extraordinary magnetic properties, including strong exchange coupling, long-range magnetic ordering, and strong magnetic anisotropy [25,26,34–37]. The other fascinating physical functionalities, such as photoinduced magnetic and magneto-optical phenomena, microporosity, catalytic activity, ferroelectricity, and ionic conductivity have been found for the cyanido-bridged coordination compounds [38–42]. Primarily, these properties were investigated within the classical group of three-dimensional bimetallic coordination frameworks based on hexacyanidometallates, called Prussian Blue Analogues (PBAs) [43]. However, the last several years brought the increasing attention pointed to the heterometallic coordination architectures based on other polycyanidometallates, including the homoligand tetracyanido-metallates of Pt\(^{II}/\)Pd\(^{II}\), heptacyanidometallates of Mo\(^{III}/\)Re\(^{IV}\), and octacyanidometallates of Mo\(^{IV}/\)W\(^{IV}/\)Re\(^{V}\), Nb\(^{IV}\),
and the heteroligand $[\text{M(L)}_x(\text{CN})_y]^n^-$ cyanide complexes with blocking organic ligand, L [44–49]. As a result, a variety of the $[\text{M(CN)}_2]^n^-$-based heterometallic coordination networks were reported, and many extraordinary magnetic, optical, and magneto-optical effects have been presented [50–54]. Some of the polycyanidometallates were also employed in the construction of luminescent functional materials, most successfully realized by combining cyanide metal complexes with lanthanide(3+) ions. Thus, in this review, we present the complete overview of luminescent heterometallic cyanido-bridged networks incorporating 4f metal ions with the emphasis on their structural features and optical properties, including sensitized and/or tunable emission. We focus and discuss only on the lanthanide-containing coordination systems based on polycyanido-metallates for which the photoluminescent properties have been recognized. This review covers the literature sources from 1970s, when this scientific area started, to September 2017.

2. Dicyanidometallates, $[\text{M}^I(\text{CN})_2]^-$ ($\text{M} = \text{Ag, Au}$)

Thanks to the $d^{10}$ valence electron configuration, dicyanidoargentate(I) and dicyanidoaurate(I) ions are almost colourless as they do not efficiently absorb the visible light. In addition, they are photoluminescent in the visible range due to their metal-to-ligand charge transfer (MLCT) transition [27,28]. They were found to be attractive for the sensitization of lanthanide(3+) luminescence, which was precisely investigated for the family of classical three-dimensional $[\text{Ln}^{III}(\text{H}_2\text{O})_3][\text{M}^I(\text{CN})_2]_3$ ($\text{Ln} = \text{La, Ce, Pr, Nd, Sm, Eu, Gd, Tb, Dy}; \text{M} = \text{Ag, Au}$) coordination polymers (Figure 1, Table 1) [55–69]. These compounds are composed of the alternately arranged $\text{Ln}^n$ and $\text{M}^I$ ($\text{M} = \text{Ag, Au}$) centres bonded through cyanide bridges (Figure 1a). Each lanthanide(3+) ion coordinates six cyanides aligned in the nearly trigonal prismatic geometry, and three additional water molecules completing the nine-coordinated metal complex. The whole network is further propagated by the almost linear $[\text{M}^I(\text{CN})_2]^-$ linkages which produce the hexagonal-type of 3D coordination network (Figure 1a) [60]. Under the UV light excitation, dicyanidometallates(I), closely connected with lanthanides, transfer the energy to the 4f metal centre resulting in the sensitized photoluminescence. The significant $[\text{M}^I(\text{CN})_2]^-$-to-$\text{Ln}^{III}$ energy transfer (ET) process was detected for the Pr-, Eu-, Tb-, and Dy-containing LnAg/LnAu networks [55,56,59,61,65]. For EuAu compound, the efficient ET effect was found under an ambient pressure, and the emission spectrum is dominated by the sharp peaks of Eu$^{3+}$ (Figure 1b). Interestingly, the increase of pressure leads to the shift of the $[\text{Au}^I(\text{CN})_2]^-$-based broad emission towards the lower energies which weakens the spectral overlap with Eu$^{3+}$, and the efficiency of the radiationless energy transfer. Therefore, with increasing pressure, the $[\text{Au}^I(\text{CN})_2]^-$-based luminescent component increases, coexisting with the Eu$^{3+}$ emission, and the complete domination of the broad dicyanidoaurate(I) emission is observed above 50 kbar [57]. The similar effect, but caused by the temperature change, was reported for TbAu analogue. In this material, the cooling procedure leads to the red shift of $[\text{Au}^I(\text{CN})_2]^-$-based emission increasing the spectral overlap with the absorption band of Tb$^{III}$ which produces the enhanced 4f-centred photoluminescence (Figure 1b) [58]. For the lanthanide(3+) ions, which do not reveal the visible emission (La, Nd, Gd), the related 3D LnAg and LnAu networks exhibit the $[\text{M}^I(\text{CN})_2]^-$-centred luminescence, also sensitive to the changes of pressure and temperature [58,62,64]. Moreover, the composition-dependent visible photo-luminescence was presented for the mixed trimetallic LaAgAu and TbAgAu compounds [62,63], the very weak energy transfer from 4d/5d to 4f metal centres was detected, resulting in the complex UV-Vis emission signal of the mixed Ln$^{3+}$ and $[\text{M}^I(\text{CN})_2]^-$ origins [66–69]. In the case of Sm$^{3+}$, even the reverse Sm(III) to Au(I) energy transfer could be postulated [69].

The combination of dicyanidoaurates(I) with lanthanide(3+) ions in the non-aqueous solution hampered the crystallization of the 3D network, and gave cyanido-bridged ($^4\text{Bu}_4\text{N})_2[\text{Ln}^{III}(\text{NO}_3)_4]$ $[\text{Au}^I(\text{CN})_2]_2$ ($\text{Ln} = \text{Ce, Nd, Sm, Gd, Eu, Tb, Dy}$) zig-zag chains (Figure 2) [67–70]. For this coordination topology, unlike to the related 3D LnAu networks, there is no efficient energy transfer between $[\text{Au}^I(\text{CN})_2]^-$ and Ln$^{III}$ complexes, and both entities act as two separate emissive sources. This was explained by the spatial separation of the 1D bimetallic LnAu coordination polymers by the $[^4\text{Bu}_4\text{N}]^+$...
cations. It causes the lack of aurophilic Au–Au interactions (Figure 2a), playing the presumably important role in the ET process within the 3D LnAu networks [67]. Therefore, taking advantage of the intrinsic properties of both luminescent 5d and 4f (Sm, Eu, Tb, Dy) metal complexes embedded in such cyanido-bridged chains, the multi-coloured composition-dependent emission, including the white light, was fruitfully achieved (Figure 2b) [67–70]. Moreover, the related trimetallic mixed 5d-4f-4f’ compounds enabled the adjustment of the emission colour to the very broad range of the visible spectrum between red colour characteristic of EuAu chains, green emission typical for TbAu chains, up to the deep violet detected for CeAu analogues, for which mainly the UV-violet CeIII-centred emission was found (Figure 2b) [68,69]. In addition, the selective excitation of the AuI-or LnIII-based emission transitions was also achievable, as exemplified by the multi-coloured photoluminescence of the mixed Ce0.33Eu0.17Tb0.5Au chains (Figure 2b) [69].

Figure 1. Crystal structure and optical properties of three-dimensional [LnIII(H2O)3][M(I)(CN)2]3 (Ln = La, Ce, Pr, Nd, Sm, Eu, Gd, Tb, Dy; M = Ag, Au) cyanido-bridged coordination polymers: (a) the views of the representative fragment of the structure together with the views of the whole network along a and c crystallographic directions, (b) the low-temperature (T = 20 K, λexc = 364 nm) pressure-dependent emission spectra of EuAu network, and the temperature-dependent emission spectra of TbAu compound (λexc = 350 nm). Colours for the structural diagrams: Ln, red; Ag/Au, dark blue; CN−, blue; H2O, grey [57,58,60]. Reprinted with permission from Inorg. Chem. 1998, 37, 3209–3316. Copyright 1998 American Chemical Society. Reprinted with permission from Inorg. Chem. 2000, 39, 4527–4534. Copyright 2000 American Chemical Society.

The further reduction of coordination dimensionality for lanthanide(III)–dicyanidometallates(I) compounds was achieved by the implementation of N,N,N-tridentate 2,2′:6′:2″-terpyridine (terpy) ligand. The resulting discrete dinuclear [LnIII(terpy)(H2O)(NO3)2][AuI(CN)2] (Ln = Sm, Eu, Gd, Tb, Dy, Ho, Er, Yb) molecules exhibit the various lanthanide-dependent emission properties [71,72]. In the case of EuAu and TbAu, the dominant visible 4f-centred luminescence was detected due to the efficient intramolecular terpy-to-Ln3+ and partial AuI-to-Ln3+ energy transfer effects. It is accompanied by the residual broad green emission related to the transitions of the [AuI(CN)2]−2 dimeric excimers that are formed between the cyanido-bridged molecules. The AuI-based emission is dominant for the other lanthanides where the ET processes to 4f metal centre are much less favoured [72]. In GdAu analogue
involving Gd$^{3+}$ ion, not luminescent in the visible range, the [Au(CN)$_2$]$^-$ based green emission is also observed but the emission spectrum is dominated by the other intense red photoluminescence, presumably assigned to the formation of an excited state exciplex of the closely stacked intermolecular terpy ligands [71].

Table 1. Photoluminescent dicyanometallate-based compounds containing lanthanide ions.

| Compound | Structural Type $^3$ | Luminescent Property | Ref. |
|----------|----------------------|----------------------|------|
| [Ln$^{III}$(NO$_3$)$_2$][Au(CN)$_2$] | 3D network | Visible Ln$^{3+}$ (Pr, Eu, Tb, Dy) emission with pressure- and temperature-dependent [M(CN)$_2$]$^-$ -to-Ln$^{3+}$ MLCT. UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). UV-Vis pressure- and temperature- tunable emission of [M(CN)$_2$]$^-$ (MLCT). | $^{55–69}$ |
| (*Bu$_4$N)$_2$[Ln$^{III}$NO$_3$][Au(CN)$_2$] | 1D zig-zag chain | Excitation- and composition- tunable visible [Au(CN)$_2$]$^-$ and Ln$^{3+}$ (Sm, Eu, Tb, Dy) emission in LnAu. UV-violet Ce$^{3+}$ emission in CeAu. | $^{67–70}$ |
| [Ln$^{III}$((terpy)(H$_2$O))(NO$_3$)$_2$][Au(CN)$_2$] | 0D dinuclear molecule | Visible Ln$^{3+}$ (Sm, Eu, Tb, Dy) and green [Au(CN)$_2$]$^-$ emission with terpy-to-Ln$^{3+}$ and partial [Au(CN)$_2$]$^-$ to-Ln$^{3+}$ ET. Mixed green [Au(CN)$_2$]$^-$ and red terpy-based emission in GdAu. | $^{71,72}$ |
| [Ln$^{III}$((bbp)(NO$_3$)$_2$)][Au(CN)$_2$]·MeCN | 0D tetranuclear molecule | Red Eu$^{3+}$ (EuAu) emission with bbp-to-Ln$^{3+}$ ET. Mixed green [Au(CN)$_2$]$^-$ and green bbp emission in GdAu. | $^{73}$ |
| [Ln$^{III}$((bbp)(NO$_3$)$_2$)][Au(CN)$_2$] | 1D linear chain | Gd Tb$^{3+}$ (GdAu) or red Eu$^{3+}$ (EuAu) emission with bbp-to-Ln$^{3+}$ ET. Green bbp-based emission in GdAu. | $^{73}$ |

$^1$ cyanido-bridged skeleton, excluding the Ag–Ag or Au–Au short contacts; $^2$ ET = energy transfer; $^3$ MLCT = metal-to-ligand charge transfer; $^4$ terpy = 2,2':6',2"-terpyridine; $^5$ bbp = bis(benzimidazole)pyridine.

Figure 2. Crystal structure and optical properties of cyanido-bridged (*Bu$_4$N)$_2$[Ln$^{III}$NO$_3$][Au(CN)$_2$] (Ln = Ce, Nd, Sm, Gd, Eu, Tb, Dy) chains: (a) the views of the single coordination chain, and the arrangement of chains, (b) the multi-coloured lanthanide-dependent emission of LnAu chains together with the related photos under UV light (top), and the multi-coloured temperature- and excitation-dependent emission of Ce$_{0.33}$Eu$_{0.17}$Tb$_{0.5}$Au chains together with the photo under UV light, all presented on the CIE 1931 chromaticity diagram. Colours for the structural diagrams: Ln, red; Au, dark blue; CN$^-$, blue; NO$_3^-$ and *Bu$_4$N$^+$, orange [67,69]. Reprinted with permission from Inorg. Chem. 2017, 56, 7948–7959. Copyright 2017 American Chemical Society.
The interesting metal-metal-ligand interactions were observed in the series of coordination systems built of dicyanidoaurate(I) ions, and lanthanide(III) complexes with N,N,N-tridentate bis(benzimidazole)pyridine (bpp) ligands [73]. Depending on the synthetic conditions, the cyanido-bridged tetranuclear \([\text{Ln}^{III}\text{(bpp)}_2][\text{Au}^{I}(\text{CN})_2]_3\cdot2\text{MeCN} (\text{Ln} = \text{Eu}, \text{Gd})\) molecules, or linear \([\text{Ln}^{III}(\text{bpp})(\text{NO}_3)_2][\text{Au}^{I}(\text{CN})_2]\cdot\text{MeCN} (\text{Ln} = \text{Eu}, \text{Gd}, \text{Tb})\) chains were obtained. For Eu- and Tb-containing species the dominant 4f-centred photoluminescence was achieved, and the efficient bpp-to-Ln\(^{3+}\) energy transfer was postulated, while the \([\text{Au}^{I}(\text{CN})_2]^-\) ions are rather optically silent in this particular case. However, due to the present Au–Au interactions observed between the tetranuclear \([\text{LnAu}_3]\) molecules, the violet \([\text{Au}^{I}(\text{CN})_2]^-\)-based emission could be detected but only for the GdAu analogue. Such 5d-centred photoluminescence was not observed for the cyanido-bridged LnAu chains lacking of the aurophilic interactions. All the related Gd-containing materials show additionally green emission of the bpp ligands. Photoluminescent dicyanidometallate-based coordination systems involving lanthanide ions are summarized in Table 1.

### 3. Tetracyanidometallates, \([\text{M}^{II}(\text{CN})_4]^{2-}\) (\(\text{M} = \text{Ni}, \text{Pd}, \text{Pt}\))

Among the square planar tetracyanidometallates(II) of group 10 metals, the tetracyanido-platinate(II) ion arouses the greatest interest in the construction of photoluminescent materials as it reveals the strong visible emission. It originates from the metal-to-metal-to-ligand charge transfer (MMLCT) transitions of the one-dimensional \([\text{Pt}^{II}(\text{CN})_4]^{2-}\) stacks controlled by the short Pt–Pt contacts [29]. As this emission is closely related to the alignment of square planar metal complexes, it was found to be strongly anisotropic, and the change of the light polarization induces the drastic change in the intensity of a few emission components assignable to the various electronic transitions. As a result, the colour of \([\text{Pt}^{II}(\text{CN})_4]^{2-}\) photoluminescence is polarization-dependent, and varies in the broad range from approximately blue to red.

This phenomenon was nicely investigated for the family of classical cyanido-bridged \([\text{Ln}^{III}(\text{H}_2\text{O})_{n+1}]_2[\text{Pt}^{II}(\text{CN})_4]_2[\text{Pt}^{II}(\text{CN})_4]\cdot9\text{H}_2\text{O} (n = 6, \text{Ln} = \text{La–Lu}; n = 5.5, \text{Ln} = \text{Eu}, \text{Tb})\) networks (Figure 3, Table 2) [74–79]. They are constructed of the coordination layers of a 6-membered metal rings topology based on Pt\(^{II}\) and Ln\(^{III}\) centres bridged by cyanide ligands within the ab plane (Figure 3a). The cyanido-bridged layers are further connected by the close Pt–Pt interactions giving the 1D \([\text{Pt}^{II}(\text{CN})_4]^{2-}\) stacks arranged along c crystallographic axis. For most of the lanthanide(3+) ions, such LnPt networks reveal mainly the strong and polarization-dependent broad emission of the \([\text{Pt}^{II}(\text{CN})_4]^{2-}\) stacks [74–77]. The emission colour and intensity were found to be dependent on the type of accompanied lanthanide(3+) ions due to the influence of the characteristic excited states of 4f metal ions amending the non-radiative processes within the bimetallic networks [75,77]. Moreover, the LuPt network exhibits the strong impact of the applied magnetic field on the Pt\(^{II}\)-based photoluminescence due to the role of 4f orbitals of Lu\(^{III}\), while the other lanthanides show only very weak emission changes upon the variable magnetic field [76].
wavelengths, the EuPt chains exhibit exclusively the sharp emission peaks assigned to Eu<sup>III</sup> which pressure. Its increase induces the red shift of the Pt<sup>II</sup>-centred photoluminescence in the visible range indicating the efficient [Pt<sup>II</sup>(CN)<sub>4</sub>]<sup>2-</sup> to-Ln<sup>3+</sup> energy transfer. The polarization-dependent Pt<sup>II</sup>-centred green to orange emission is caused by the increase of temperature, clearly hampering the lanthanide photoluminescence [79]. The similar effect is caused by the increase of temperature, clearly hampering the lanthanide photoluminescence [79]. The strong influence of the Pt–Pt distances on the optical properties was proved by comparison of the closely related [Er<sup>III</sup>(H<sub>2</sub>O)<sub>6</sub>]<sup>2-</sup> ions to 4f metal centre (Figure 4b) [81]. On the contrary, the TbPt chains are still observed but significantly decreased through the radiationless energy transfer. The sensitization process towards Sm<sup>3+</sup> and Eu<sup>3+</sup> was reported to be sensitive to the hydrostatic pressure. Its increase induces the red shift of the Pt<sup>II</sup>-centred emission reducing the spectral overlap with Ln<sup>3+</sup> absorption states that leads to the gradual disappearance of the 4f-centred emission [78,79]. The similar effect is caused by the increase of temperature, clearly hampering the lanthanide photoluminescence [79]. The strong influence of the Pt–Pt distances on the optical properties was proved by comparison of the closely related [Er<sup>III</sup>(H<sub>2</sub>O)<sub>6</sub>]<sup>2-</sup> ions to 4f metal centre (Figure 4b) [81]. On the contrary, the TbPt chains are still observed but significantly decreased through the radiationless energy transfer. The sensitization process towards Sm<sup>3+</sup> and Eu<sup>3+</sup> was reported to be sensitive to the hydrostatic pressure. Its increase induces the red shift of the Pt<sup>II</sup>-centred emission reducing the spectral overlap with Ln<sup>3+</sup> absorption states that leads to the gradual disappearance of the 4f-centred emission [78,79]. The similar effect is caused by the increase of temperature, clearly hampering the lanthanide photoluminescence [79]. The strong influence of the Pt–Pt distances on the optical properties was proved by comparison of the closely related [Er<sup>III</sup>(H<sub>2</sub>O)<sub>6</sub>]<sup>2-</sup> ions to 4f metal centre (Figure 4b) [81]. On the contrary, the TbPt chains are still observed but significantly decreased through the radiationless energy transfer. The sensitization process towards Sm<sup>3+</sup> and Eu<sup>3+</sup> was reported to be sensitive to the hydrostatic pressure. Its increase induces the red shift of the Pt<sup>II</sup>-centred emission reducing the spectral overlap with Ln<sup>3+</sup> absorption states that leads to the gradual disappearance of the 4f-centred emission [78,79]. The similar effect is caused by the increase of temperature, clearly hampering the lanthanide photoluminescence [79]. The strong influence of the Pt–Pt distances on the optical properties was proved by comparison of the closely related [Er<sup>III</sup>(H<sub>2</sub>O)<sub>6</sub>]<sup>2-</sup> ions to 4f metal centre (Figure 4b) [81]. On the contrary, the TbPt chains are still observed but significantly decreased through the radiationless energy transfer. The sensitization process towards Sm<sup>3+</sup> and Eu<sup>3+</sup> was reported to be sensitive to the hydrostatic pressure. Its increase induces the red shift of the Pt<sup>II</sup>-centred emission reducing the spectral overlap with Ln<sup>3+</sup> absorption states that leads to the gradual disappearance of the 4f-centred emission [78,79]. The similar effect is caused by the increase of temperature, clearly hampering the lanthanide photoluminescence [79].

In contrast to the other 4f metal ions, SmPt and EuPt networks reveal the strong lanthanide photoluminescence in the visible range indicating the efficient [Pt<sup>II</sup>(CN)<sub>4</sub>]<sup>2-</sup> to-Ln<sup>3+</sup> energy transfer mechanism (Figure 3b) [75,77–79]. The polarization-dependent Pt<sup>II</sup>-based green to orange emission is caused by the increase of temperature, clearly hampering the lanthanide photoluminescence [79]. The similar effect is caused by the increase of temperature, clearly hampering the lanthanide photoluminescence [79]. The strong influence of the Pt–Pt distances on the optical properties was proved by comparison of the closely related [Er<sup>III</sup>(H<sub>2</sub>O)<sub>6</sub>]<sup>2-</sup> ions to 4f metal centre (Figure 4b) [81]. On the contrary, the TbPt chains are still observed but significantly decreased through the radiationless energy transfer. The sensitization process towards Sm<sup>3+</sup> and Eu<sup>3+</sup> was reported to be sensitive to the hydrostatic pressure. Its increase induces the red shift of the Pt<sup>II</sup>-centred emission reducing the spectral overlap with Ln<sup>3+</sup> absorption states that leads to the gradual disappearance of the 4f-centred emission [78,79]. The similar effect is caused by the increase of temperature, clearly hampering the lanthanide photoluminescence [79]. The strong influence of the Pt–Pt distances on the optical properties was proved by comparison of the closely related [Er<sup>III</sup>(H<sub>2</sub>O)<sub>6</sub>]<sup>2-</sup> ions to 4f metal centre (Figure 4b) [81]. On the contrary, the TbPt chains are still observed but significantly decreased through the radiationless energy transfer. The sensitization process towards Sm<sup>3+</sup> and Eu<sup>3+</sup> was reported to be sensitive to the hydrostatic pressure. Its increase induces the red shift of the Pt<sup>II</sup>-centred emission reducing the spectral overlap with Ln<sup>3+</sup> absorption states that leads to the gradual disappearance of the 4f-centred emission [78,79]. The similar effect is caused by the increase of temperature, clearly hampering the lanthanide photoluminescence [79].

Following the promising optical properties of 3D Ln<sup>III</sup>–[Pt<sup>II</sup>(CN)<sub>4</sub>]<sup>2-</sup> networks, a considerable number of the related d–f cyanido-bridged coordination polymers revealing photoluminescent functionalities were reported (Table 2) [80–88]. The insertion of 2,2′:6′:2″-terpyridine (terpy) into the self-assembled Ln<sup>III</sup>–Pt<sup>II</sup> system resulted in the one-dimensional [Ln<sup>III</sup>(terpy)(H<sub>2</sub>O)<sub>2</sub>(NO<sub>3</sub>)][Pt<sup>II</sup>(CN)<sub>4</sub>]<sup>2-</sup>·n(solvent) (Ln = Eu, Tb) zig-zag chains (Figure 4a) [81,87]. Under UV light excitation of various wavelengths, the EuPt chains exhibit exclusively the sharp emission peaks assigned to Eu<sup>III</sup> which...
clearly indicates the efficient dual acceptor–donor intramolecular energy transfer from both terpy ligand and \([\text{Pt}^{II}(\text{CN})_4]^{2-}\) ions to 4f metal centre (Figure 4b) \[81\]. On the contrary, the TbPt chains shows the dual visible photoluminescence, the green 4f-centred and the blue \([\text{Pt}^{II}(\text{CN})_4]^{2-}\) based emission. Thus, only the terpy-to-Tb\(^{3+}\) energy transfer is efficient while the analogous Pt\(^{II}\)-to-Tb\(^{3+}\) process is partial. It leaves two distinguishable emission sources that can be selectively induced, leaving the possibility to observe the separate 4f- and ligand-centred photoluminescence. Such a Pt–Pt contacts. Thus, it is weakened for the compounds lacking of such platinophilic interactions, leaving the possibility to observe the separate 4f- and ligand-centred photoluminescence. Such a

The MMLCT emission of \([\text{Pt}^{II}(\text{CN})_4]^{2-}\) ions is strongly determined by the presence of short Pt–Pt contacts. Thus, it is weakened for the compounds lacking of such platinophilic interactions, leaving the possibility to observe the separate 4f- and ligand-centred photoluminescence. Such a
situation was found in several tetracyanidometallate-based d–f coordination systems with various organic ligands (Table 2) [80,83–85]. For instance, the Ln³⁺ (Ln = Er, Yb) complexes with tetraphenylporphinirinate dianion combined with various tetracyanidometallates resulted in the trinuclear [LnIII(tpp)(dmf)(H₂O)₂][M(II)(CN)₄] (Ln = Er, Yb; M = Ni, Pd, Pt) molecules, for which the exclusively red tpp-based emission along with the characteristic Er³⁺ and Yb³⁺ NIR phosphorescence peaks were observed [80]. Similar effects of 4f-centred emission realized by a simple direct f–f excitation or organic ligand-to-Ln³⁺ energy transfer were detected for a series of the ionic salts of lanthanide(3+) ions with dimethylsulfoxide, 1,10-phenanthroline (phen) and 2,2'-bipyridine accompanied by non-coordinated tetracyanidometallates, and for the trinuclear [LnIII₂PdIII] molecules with the supporting phen ligand [84,85]. Only in the [EuIII(dmfo)(terpy)(H₂O)₂(NO₃)]·[Pt(II)(CN)₄] ionic salt without platinophilic interactions, the weak PtIV-based visible emission along with the stronger terpy-sensitized EuIII photoluminescence could be detected [83]. On the other hand, two-dimensional K[LnIII(H₂O)₃][Pt(II)(CN)₄]·[Pt(II)(CN)₄]·5.5H₂O (Ln = La, Pr, Nd) honeycomb networks bearing the numerous [Pt(II)(CN)₄]²⁻ stacks gives the PtII-based green emission, very strong for LaPt, and partially impaired by the excited states of PrIII and NdIII in PrPt and NdPt, respectively [88].

Table 2. Photoluminescent tetracyanidometallate-based compounds containing lanthanide ions.

| Compound | Structural Type | Luminescent Property | Ref. |
|----------|----------------|----------------------|------|
| [LnIII(H₂O)₃][Pt(II)(CN)₄]·[Pt(II)(CN)₄]·9H₂O (Ln = La–Lu) | 2D 6-membered metal rings | Visible polarization-dependent emission from 1D [Pt(CN)₄]²⁻ stacks (MMLCT). | [74–76] |
| [LnIII₂(H₂O)₃][Pt(II)(CN)₄]·2Pt(II)(CN)₄·10H₂O (Ln = Sm, Eu) | 2D 6-membered metal rings | Polarization-dependent green to orange emission from 1D [Pt(CN)₄]²⁻ stacks. Red Eu³⁺ (EuPt) or Sm³⁺ (SmPt) emission with temperature- and pressure-dependent [Pt(CN)₄]²⁻ to-LnIII ET. | [74,77–79] |
| [ErIII(H₂O)(SO₄)][Pt(II)(CN)₄]·[Pt(II)(CN)₄]·1.5H₂O | 2D 6-membered metal rings | Polarization-dependent green to orange emission from 1D [Pt(CN)₄]²⁻ stacks. Red tpp, and NIR Er³⁺ (ErM) or Yb³⁺ (YbM) emission. | [74] |
| [LnIII(tpp)(dmf)(H₂O)₂][M(II)(CN)₄] | 0D trinuclear molecule | Red Eu³⁺ emission with terpy-to-Eu³⁺ and [Pt(CN)₄]²⁻ to-Eu³⁺ ET. | [81,82] |
| [EuIII(terpy)(H₂O)₂][Pt(II)(CN)₄]·MeCN | 1D zig-zag chain | Red Eu³⁺ emission with terpy-to-Eu³⁺ and [Pt(CN)₄]²⁻ to-Eu³⁺ ET. | [82] |
| [EuIII(terpy)(H₂O)₃][Pt(CN)₄]·2H₂O | 1D ladder chain | Red Eu³⁺ emission with terpy-to-Eu³⁺ and [Pt(CN)₄]²⁻ to-Eu³⁺ ET. | [82] |
| [EuIII(terpy)(H₂O)₃][Pt(CN)₄]·3H₂O | 0D ionic salt | Red Eu³⁺ emission with terpy-to-Eu³⁺ ET, additional emission from [Pt(CN)₄]²⁻. | [83] |
| [EuIII(dmfo)(terpy)(H₂O)₂][Pt(II)(CN)₄] | 0D ionic salt | Red Eu³⁺ emission with terpy-to-Eu³⁺ ET, additional emission from [Pt(CN)₄]²⁻. | [83] |
| [LnIII(dmfo)(H₂O)₃][Pt(II)(CN)₄]·0.5[M(II)(CN)₄]·2H₂O | 0D ionic salt | Red Eu³⁺ (EuM) or green Tb³⁺ (TbM) emission with direct f–f excitation. | [84] |
| [LnIII(dmfo)(phen)(H₂O)₂][Pt(II)(CN)₄] | 0D trinuclear molecule | Red Eu³⁺ (EuPt) or green Tb³⁺ (TbPt) emission with phen-to-LnIII ET. | [85] |
| [LnIII(dmfo)(phen)(H₂O)₂][Pt(II)(CN)₄] | 0D trinuclear molecule | Red Eu³⁺ (EuPt) or green Tb³⁺ (TbPt) emission with phen-to-LnIII ET. | [85] |
| [LnIII(dmfo)(2,2'-bpy)(H₂O)₂][Pt(II)(CN)₄] | 0D ionic salt | Red Eu³⁺ (EuPt) or green Tb³⁺ (TbPt) emission with 2,2'-bpy-to-LnIII ET. | [85] |
| K₂[TbIII(H₂O)₃][Pt(II)(CN)₄]·[Au(III)(CN)₂]·2H₂O | 2D 6-membered metal rings | Green Tb³⁺ emission with [Au(III)₂]-to-Tb³⁺ ET; additional weak blue [Au(III)₂] emission. | [86] |
| [TbIII(2,2'-bpy)(H₂O)₃][Pt(II)(CN)₄]·[Au(III)(CN)₂]·1.5[2,2'-bpy]·2H₂O | 0D ionic salt | Green Tb³⁺ emission with [Au(III)₂]-to-Tb³⁺ ET; additional weak violet 2,2'-bpy emission. | [87] |
| [TbIII(terpy)(ClO₄)(NO₃)][Pt(II)(CN)₄]·[Cu(II)(CN)₄]·2.5H₂O·10H₂O | 1D zig-zag chain | Green Tb³⁺ emission with terpy-to-Tb³⁺ ET; additional blue [Pt(CN)₄]²⁻ emission. | [87] |
| [TbIII(terpy)(H₂O)₂][ClO₄][Pt(II)(CN)₄]·[Cu(II)(CN)₄]·4H₂O | 0D ionic salt | Green Tb³⁺ emission with terpy-to-Tb³⁺ and CH₃COO⁻·to-Tb³⁺ ET. | [87] |
| [TbIII(terpy)(H₂O)₂][ClO₄][Pt(II)(CN)₄]·[Cu(II)(CN)₄]·7H₂O | 0D ionic salt | Green Tb³⁺ emission with terpy-to-Tb³⁺ and CH₃COO⁻·to-Tb³⁺ ET. | [87] |
| K[LnIII(H₂O)₃][Pt(II)(CN)₄]·[Pt(II)(CN)₄]·5.5H₂O (Ln = La, Pr, Nd) | 2D honeycomb | Green emission from 1D [Pt(CN)₄]²⁻ stacks (MMLCT). | [88] |

1. cyanido-bridged skeleton, excluding the Pt–Pt and the analogous M–M short contacts; 2. MMLCT = metal-metal-to-ligand charge transfer; 3. ET = energy transfer; 4. terpy = 2,2'6,2''-terpyridine; 5. tpp = tetraphenylporphinirinate dianion; 6. dmfo = N,N,N'-dimethylformamide; 7. dms = dimethylsulfoxide; 8. phen = 1,10-phenanthroline; 9. 2,2'-bpy = 2,2'-bipyridine; 10. terpyCl = 4'-chloro-2,2',6,2''-terpyridine.

Tetracyanidoplatinate(II) together with dicyanodaurate(II) were also employed in construction of trimetallic TbIII–PdIII–AuI coordination networks [86]. The resulting K₂[TbIII(H₂O)₃][Pt(II)(CN)₄]·[Au(III)(CN)₂]·2H₂O layers are composed of TbIII bridged by cyanides to PtII which produces the 2D
coordination polymer of an 8-membered metal rings topology. The non-cyanido-bridging $\text{[Au}^\text{(CN)}_2]^{2-}$ moieties are stacked within one type of metal rings, and interact with $\text{[Pt}^\text{(CN)}_4]\text{[Au}^\text{(CN)}_2]^{2-}$ by Pt–Au and Au–Au short contacts. This results in the formation of $\text{[Au}_3\text{Pt}_4]$ metallophilic clusters of their own blue emission. Moreover, these clusters are able to sensitize green Tb$^{3+}$ emission. Upon the addition of 2,2’-bipyridine (2,2’-bpy), the cyanido-bridged heterotrinuclear $\text{[Tb}^\text{III}(2,2’\text{-bpy})(\text{H}_2\text{O})_4]\text{[Pt}^\text{III}(\text{CN})_4]\text{[Au}^\text{(CN)}_2]^{-1.5(2,2’-bpy)}\text{2H}_2\text{O}$ molecules were achieved. The $\text{[Tb}^\text{III}\text{Pt}_2\text{Au}]$ molecules are also aggregated through Au–Au and Au–Pt interactions, and the tetrameric $\text{[Au}_2\text{Pt}_2]$ metallophilic clusters were detected. This material reveals the green Tb$^{3+}$ emission with $\text{[Au}_2\text{Pt}_2]$-to-Tb$^{3+}$ energy transfer, and the additional violet 2,2’-bpy emission, proving that $\text{[Au}^\text{(CN)}_2]^{-1},\text{[Pt}^\text{(CN)}_4]^{2-}$, and their metallophilic clusters can serve as good sensitizers for the lanthanide(3+) photoluminescence [86].

4. Hexacyanidometallates, $\text{[M}^\text{III}(\text{CN})_n]^3-$ ($\text{M = Cr, Co}$)

Among the rich diversity of reported hexacyanidometallates [43], only hexacyanidocobaltate(III) and hexacyanidocobaltate(III) ions were applied in the construction of photoluminescent d-f coordination networks (Figures 5 and 6, Table 3) [89–100]. They reveal the near-infrared and red emission bands, respectively, due to their metal-centred d-d electronic transitions [30,31]. In combination with trivalent lanthanide ions, they form the three-dimensional cyanido-bridged $\text{[Ln}^\text{III}(\text{H}_2\text{O})_n]\text{[M}^\text{III}(\text{CN})_n]\text{2H}_2\text{O}$ ($\text{M = Cr, Co; } n = 3$, Ln = La–Nd; $n = 2$, Nd–Lu) networks with all cyanides bridging to the neighbouring 9- or 8-coordinated 4f metal centres [89,90]. The GdCo and GdCr networks exhibit the d-metal-centred characteristic photoluminescence, observed even at room temperature due to the heavy atom effect [91]. Among the other analogues, the emission properties of the EuCo network was only reported, and red Eu$^{3+}$ photoluminescence sensitized by the Co$^{3+}$-to-Eu$^{3+}$ energy transfer was presented [92,93]. It suggested that $\text{[Co}^\text{III}(\text{CN})_6]^{3-}$ can be a reasonable sensitizer for the visible photoluminescence of 4f metal ions.

![Figure 5. Crystal structure and optical properties of $\text{[Ln}^\text{III}(\text{3-OHpy})_2(\text{H}_2\text{O})_4]\text{[Co}^\text{III}(\text{CN})_6]\text{H}_2\text{O}$ (Ln = Eu, Tb, Dy; 3-OHpy = 3-hydroxypyridine) cyanido-bridged chains: (a) the views of the coordination chain, and the arrangement of chains; (b) the white light emission of DyCo chains and the related photo (top), the multi-coloured composition-dependent emission of Eu$_x$Tb$_{1-x}$Co (bottom), all shown on the CIE 1931 chromaticity diagram. Colours for the structural diagram: Ln, red; W, dark blue; CN$^{-}$, blue; 3-OHpy, orange; H$_2$O, grey [96,97].](image-url)
Hexacyanidometallates of Co$^{3+}$ and Cr$^{3+}$ exhibit their own emission close to the edge between visible and NIR ranges, so they were expected to be good sensitizers for NIR-emitting lanthanide($3^+$) ions. It was accurately proved for the cyanido-bridged [Ln$^{III}$I(dmfi)$_4$(H$_2$O)$_2$][Cr$^{III}$I(CN)$_6$] · nH$_2$O (Ln = Nd, Gd, Yb) chains, and dinuclear [Ln$^{III}$I(dmfi)$_4$(H$_2$O)$_3$][Co$^{III}$I(CN)$_6$] · nH$_2$O (Ln = Nd, Gd, Yb) molecules [94,95]. The NdCr, NdCo, YbCr, and YbCo materials show the NIR emission of Nd$^{3+}$ or Yb$^{3+}$, and the lack of the expected d-metal-centred luminescence which indicates the efficient Cr$^{3+}$-to-Ln$^{3+}$ and Co$^{3+}$-to-Ln$^{3+}$ intramolecular energy transfer processes.

The first reports on photoluminescent hexacyanido-bridged d–f materials were expanded by the implementation of various pyrimidine and pyridine derivatives that control the coordination topology [96–100]. The combination of 3-hydroxypyridine (3-OHpy) with Dy$^{III}$ and [Co$^{III}$I(CN)$_6$]$^{3–}$ produced the cyanido-bridged [Dy$^{III}$I(3-OHpy)$_2$(H$_2$O)$_4$][Co$^{III}$I(CN)$_6$] · H$_2$O zig-zag chains (Figure 5a), exhibiting the room temperature white-light Dy$^{III}$ emission realized by the complex excitation involving not only direct f–f transitions, but also Co$^{3+}$-to-Dy$^{3+}$ and 3-OHpy-to-Dy$^{3+}$ energy transfer pathways (Figure 5b) [96]. This topology was also reported for trimetallic [Eu$^{III}$, Tb$^{III}$, 1–x (3-OHpy)$_2$(H$_2$O)$_4$][Co$^{III}$I(CN)$_6$] · H$_2$O materials showing the multi-coloured 4f-centred emission. The colour of luminescence is tuned between red, orange and yellow to green, by the compound’s composition, that is the Eu/Tb ratio governing the intensity of red Eu$^{3+}$ and green Tb$^{3+}$ emission components (Figure 5b) [97]. Multi-coloured emission is also realized for the trimetallic EuTbCo chains by the selection of the appropriate wavelengths of UV light controlling the intensities of Eu$^{3+}$ and Tb$^{3+}$ emission characteristics. Moreover, the 4f-centred photoluminescence is, here, enhanced by the energy transfer occurring from both [Co$^{III}$I(CN)$_6$]$^{3–}$ and 3-OHpy to lanthanide($3^+$) ions, and the efficiencies of these radiationless effects play a vital role in the observed switchable emission.

Figure 6. Crystal structure and optical properties of near-infrared (NIR)-emitting cyanido-bridged [Nd$^{III}$I(pmmo)$_2$(H$_2$O)$_3$][Cr$^{III}$I(CN)$_6$] (a) (pmmo = pyrimidine N-oxide) and [Yb$^{III}$I(3-pyone)$_2$(H$_2$O)$_3$][Co$^{III}$I(CN)$_6$] (b) (3-pyone = 3-pyridone) coordination polymers. The top part of the figure shows the respective coordination skeletons of square grid layers (a) and nearly linear chains (b), while the bottom part presents the room temperature excitation and NIR-emission spectra under the indicated wavelength conditions. Colours for the structural diagrams: Nd/Yb, red; Co/Cr, dark blue; CN$^–$, blue; pmmo/3-pyone, orange; H$_2$O, grey [99,100].
The replacement of 3-hydroxypyridine by 4-hydroxypyridine (4-OHpy) in the Dy\( ^{III}\)–[Co\( ^{III}\)(CN)\(_6\)]\(^{3–}\) system resulted in the dramatic change of the structure towards the cyanido-bridged \([\text{Dy}^{III}(4\text{-OHpy})\text{H}_2\text{O})\] \([\text{Co}^{III}(\text{CN})\text{H}_6]\) \(0.5\text{H}_2\text{O}\) layered framework of a 6-membered metal rings topology. It exhibits the yellow Dy\( ^{III}\) emission sensitized by the [Co\( ^{III}\)(CN)\(_6\)]\(^{3–}\) complex through intramolecular energy transfer. However, in contrast to 3-OHpy, the 4-OHpy ligand is not suitable sensitizer for Dy\( ^{III}\), and its intrinsic greenish-blue phosphorescence was detected. As a result, the multi-coloured yellow to greenish-blue emission switchable by excitation light, governing the ligand- and Dy\(^{3+}\)-based luminescent components, was achieved [98]. The layered network but of a square grid topology was prepared by the application of pyrimidine N-oxide (pmmo) with the Nd\(^{3+}\) and [Cr\( ^{III}\)(CN)\(_6\)]\(^{3–}\) ions (Figure 6a) [99]. The resulting [Nd\( ^{III}\)(pmmo)\text{H}_2\text{O}][\text{Cr}^{III}(\text{CN})\text{H}_6] material reveals the strong NIR Nd\(^{3+}\)-centred emission under the UV excitation. The observed excitation spectrum is dominated by the broad complex band assigned to the absorption of pmmo and [Cr\( ^{III}\)(CN)\(_6\)]\(^{3–}\) (Figure 6a). This, together with the disappearance of the expected green phosphorescence of organic ligand, and NIR emission of a cyanide complex, proved the efficient energy transfer from both organic and inorganic building blocks. The similar dual donor intramolecular energy transfer related to the direct coordination of the organic ligand and hexacyanidometallate to 4f metal was presented for almost linear [Yb\( ^{3+}\)(3-pyone)\text{H}_2\text{O}][\text{Co}^{III}(\text{CN})\text{H}_6] (3-pyone = 3-pyridone) chains (Figure 6b) [100]. Under the UV excitation, they show typical NIR Yb\(^{3+}\) emission, enhanced by the radiationless energy transfer from 3-pyone and [Co\( ^{III}\)(CN)\(_6\)]\(^{3–}\) to 4f metal, which proves that hexacyanidocobaltate(III) is particularly attractive photoluminescent metalloligand being a good sensitizer for both visible light- and NIR-emitting lanthanide(3+) ions.

### Table 3. Photoluminescent hexacyanidometallate-based compounds containing lanthanide ions.

| Compound | Structural Type | Luminescent Property | Ref. |
|----------|----------------|----------------------|------|
| \([\text{Gd}^{III}\text{H}_2\text{O}][\text{Co}^{III}(\text{CN})\text{H}_6]\times\text{H}_2\text{O}\) (M = Cr, Co) | 3D network | Red Co\(^{3+}\)(GdCo) or NIR Cr\(^{3+}\) (GdCr) emission | [89–91] |
| \([\text{Eu}^{III}\text{H}_2\text{O}][\text{Co}^{III}(\text{CN})\text{H}_6]\times\text{H}_2\text{O}\) (L = Nd, Gd, Yb) | 3D network | Red Eu\(^{3+}\) emission with Co\(^{3+}\)-to-Eu\(^{3+}\) ET \(^1\) | [92–93] |
| \([\text{Ln}^{III}(\text{dmf})\text{H}_2\text{O}][\text{Co}^{III}(\text{CN})\text{H}_6]\times\text{H}_2\text{O}\) (L = Nd, Gd, Yb) | 0D dinuclear molecule | NIR Nd\(^{3+}\) emission (GdCo). | [94,95] |
| \([\text{Dy}^{III}(4\text{-OHpy})\text{H}_2\text{O}][\text{Co}^{III}(\text{CN})\text{H}_6]\times\text{H}_2\text{O}\) | 1D zig-zag chain | White light Dy\(^{3+}\) emission with direct f-f excitation, Co\(^{3+}\)-to-Dy\(^{3+}\) and 3-OHpy-to-Dy\(^{3+}\) ET. | [96] |
| \([\text{Eu}^{III},\text{Th}^{IV}(3\text{-OHpy})\text{H}_2\text{O}][\text{Co}^{III}(\text{CN})\text{H}_6]\times\text{H}_2\text{O}\) | 1D zig-zag chain | Excitation and Eu/Tb ratio switching from red Eu\(^{3+}\) to green Tb\(^{3+}\) emission with Co\(^{3+}\)-to-La\(^{3+}\) ET. | [97] |
| \([\text{Dy}^{III}(4\text{-OHpy})\text{H}_2\text{O}][\text{Co}^{III}(\text{CN})\text{H}_6]\times\text{H}_2\text{O}\times\text{H}_2\text{O}\) | 2D 6-membered metal rings | Excitation switching from yellow Dy\(^{3+}\) emission with Co\(^{3+}\)-to-Dy\(^{3+}\) ET to greenish-blue-4-OHpy ET. | [98] |
| \([\text{Nd}^{III}(\text{pmmo})\text{H}_2\text{O}][\text{Co}^{III}(\text{CN})\text{H}_6]\times\text{H}_2\text{O}\) | 2D square grid | NIR Nd\(^{3+}\) emission with Co\(^{3+}\)-to-Nd\(^{3+}\) and pmmo-to-Nd\(^{3+}\) ET. | [99] |
| \([\text{Yb}^{III}(3\text{-pyone})\text{H}_2\text{O}][\text{Co}^{III}(\text{CN})\text{H}_6]\times\text{H}_2\text{O}\) | 1D chain | NIR Yb\(^{3+}\) emission with Co\(^{3+}\)-to-Yb\(^{3+}\) and 3-pyone-to-Yb\(^{3+}\) ET. | [100] |

\(^1\) ET = energy transfer; \(^2\) dmf = N,N-dimethylformamide; \(^3\) 3-OHpy = 3-hydroxypyridine; \(^4\) 4-OHpy = 4-hydroxypyridine; \(^5\) pmmo = pyrimidine N-oxide; \(^6\) 3-pyone = 3-pyridone.

### 5. Octacyanidometallates, \([\text{MO}^{IV/IV}(\text{CN})_8]\)\(^{4–/3–}\) (M = Mo, W)

Octacyanidometallate complexes of Mo(IV,V) and W(IV,V) are not emissive as their numerous d-d and charge transfer (ligand-to-metal or metal-ligand) electronic transitions were found to be rather photoreactive which was utilized in the construction of photomagnetic materials [23]. These cyanide complexes offer, however, very light yellow colour as their absorption bands are shifted to the UV range. It enables the observation of visible and NIR photoluminescence of accompanying emissive chromophores [32]. In effect, a considerable number of photoluminescent d-f coordination networks based on octacyanidometallates were reported (Figures 7 and 8, Table 4) [101–110]. In the
aqueous solution, trivalent lanthanide(3+) and [M\(^{IV}(CN)_{8}\)]\(^{3-}\) ions produce the cyanido-bridged [Ln\(^{III}(H_{2}O)_{3}][M^{V}(CN)_{6}]\) (Ln = Sm, Eu, Gd, Tb; M = Mo, W) layered frameworks of a square grid topology [101–103]. Such EuMo and EuW layers exhibit the specific Eu\(^{3+}\) emission realized partially by direct f-f excitation, and the more sophisticated energy transfer from O-Eu (EuMo) or O-W (EuW) ligand-to-metal charge transfer (LMCT) bands situated in the UV range [101]. In the analogous TbMo and TbW compounds, the green emission of Tb\(^{3+}\) was detected after the excitation by the UV light, related to the Tb-based d–f and f–f electronic transitions [101,102]. Similar Eu\(^{3+}\) or Tb\(^{3+}\) emission was reported for the mixed [Eu\(^{II}III0.5\)Gd\(^{II}0.5\)(H\(_{2}\)O)\(_{5}\)]\([W^{V}(CN)_{8}]\) and [Sm\(^{III}III0.5\)Tb\(^{III}0.5\)(H\(_{2}\)O)\(_{5}\)]\([W^{V}(CN)_{8}]\) networks. The trimetallic [Eu\(^{II}III0.5\)Tb\(^{III}0.5\)(H\(_{2}\)O)\(_{5}\)]\([W^{V}(CN)_{8}]\) framework exhibits both red Eu\(^{3+}\) and green Tb\(^{3+}\) emission peaks realized by the LMCT, O-Eu or W-related, excitation, and the additional Tb\(^{3+}\)-to-Eu\(^{3+}\) intermetallic energy transfer occurring through the [W\(^{V}(CN)_{8}\)]\(^{3-}\) moieties [103]. The emission of Nd\(^{3+}\) was also possible to detect within the family of octacyanidometallate-based materials, as reported for the cyanido-bridged [Nd\(^{III}(phen)(dmf)_{2}(H_{2}O)]\[Mo^{V}(CN)_{8}]\ 2H_{2}O \text{ and } [Nd^{III}(phen)(dmf)_{2}[M^{V}(CN)_{6}]\cdot xH_{2}O (M = Mo, W; phen = 1,10-phenanthroline) chains showing NIR emission of 4f metal ion sensitized by phen ligand [104].

The interesting luminescent functionalites were presented for the three-dimensional hybrid inorganic-organic [Ln\(^{III}(mpca)_{2}(MeOH)_{2}(H_{2}O)_{6}]\[Mo^{IV}(CN)_{8}]\cdot xMeOH (Ln = Nd, Eu, Tb; mpca = 5-methyl-2-pyrazine carboxylic acid) networks where cyanido-bridged layers are connected in the third direction by organic mpca linkers [105]. They reveal the strong 4f-metal centred photoluminescence sensitized by mpca ligands while [Mo\(^{IV}(CN)_{8}\)]\(^{4-}\) serves as an inorganic linker stabilizing the 3D coordination network. Moreover, the Eu\(^{3+}\) emission was found to be strongly sensitive to the amount of water molecules occupying the interstitial space within the crystal structure. Therefore, the emission intensity of the EuMo network increases significantly with decreasing humidity of the air around the solid sample, which makes this material an efficient humidity sensor working in the whole relative humidity range from 0 to 100% [106].
Figure 8. Crystal structure and optical properties of two-dimensional [TbIII(box)2(dmf)2] [WV(CN)8]·H2O (box = 2,2'-bis(2-oxazoline)) coordination polymers: (a) the views of the single cyanido-bridged layer, and the arrangement of layers, (b) the excitation-dependent emission spectra collected at T = 77 K. Colours for the structural diagram: Tb, red; W, dark blue; CN−, blue; box, orange; dmf, grey [109].

Table 4. Photoluminescent octacyanidometallate-based compounds containing lanthanide ions.

| Compound                              | Structural type          | Luminescent property                                      | Ref.     |
|---------------------------------------|--------------------------|-----------------------------------------------------------|----------|
| [EuIII(H2O)3][MoV(CN)8]·H2O          | 3D hybrid P2O3 network   | Red Eu3+ emission with O-Eu (EuMo) or O-W (EuW) LMCT and direct f-f excitation. | [101]    |
| TbIII(H2O)3][MoV(CN)8]·H2O          | 1D helical chain         | Thermal switching between red Eu3+ and blue Pr-pybox emission. | [102]    |
| [EuIII(phen)(dmf)3][MoV(CN)8]·H2O   | 2D square grid           | Red Eu3+ emission with phen-to-Nd3+ ET.                    | [103]    |
| TbIII(phen)(dmf)3][MoV(CN)8]·H2O   | 2D square grid           | NIR Nd3+ emission with direct f-f excitation.              | [104]    |
| Pr-pybox = (SS)-2,6-bis[8H-| 2D mixed 4- and 8-membered metal rings | Excitation switching between green Tb3+ and red box emission. | [105]    |
| LnIII(box)2(dmf)2][MoV(CN)8]·H2O  | 3,4] 1D chain            | NIR Nd3+ emission with mpca-to-Ln3+ ET, and humidity-dependent intensity of Eu3+ emission. | [106]    |
| [EuIII(box)2(dmf)2][MoV(CN)8]·H2O | 2D mixed 4- and 8-membered metal rings | Visible Ln3+ (Pr, Sm, Eu, Ho) or NIR Ln3+ (Pr, Nd, Sm, Ho, Yb) emission with direct f-f excitation and box-to-Ln3+ ET. | [106]    |
| [EuIII(box)2(dmf)2][MoV(CN)8]·H2O | 2D mixed 4- and 8-membered metal rings | Visible Ln3+ (Pr, Sm, Eu, Ho) or NIR Ln3+ (Pr, Nd, Sm, Ho, Yb) emission with direct f-f excitation and box-to-Ln3+ ET. | [106]    |

1 LMCT = ligand-to-metal charge transfer; 2 ET = energy transfer; 3 phen = 1,10-phenanthroline; 4 dmf = N,N-dimethylformamide; 5 mpca = 5-methyl-2-pyrazine carboxylic acid; 6 Pr-pybox = (SS)- or (RR)-2,2’-(2,6-pyridinediethyl)[bis(4-isopropyl-2-oxazoline)]; 7 ind-pybox = (SRSR)- or (RRS)-2,6-bis[8H-indeno[1,2-d]oxazolin-2-yl]pyridine; 8 box = 2,2’-bis(2-oxazoline).

Note: The LMCT = ligand-to-metal charge transfer; ET = energy transfer.
The implementation of the enantiopure derivatives of 2,2’-(2,6-pyridinediyldi)bis(2-oxazoline) (pybox) molecule into the LnIII-[W(V)(CN)8]3− systems resulted in the formation of chiral cyanido-bridged [LnIII(pybox)(dmf)2][W(V)(CN)8]·dmf·4H2O (Ln = Nd, Eu, Gd; pybox = (SS)- or (RR)-2,2’-(2,6-pyridinediyldi)bis(4-isopropyl-2-oxazoline); Figure 7a) and [LnIII(ind-pybox)(dmf)4][W(V)(CN)8]·5MeCN·4MeOH (Ln = Nd, Gd; ind-pybox = (SRSR)- or (RSRS)-2,6-bis[8H-indeno[1,2-d]oxazolin-2-yl]pyridine) helices [107,108]. Taking advantage of both luminescent LnIII and pybox ligand, the Eu(pybox)W chains reveal the thermal switching between red EuIII emission predominant at low temperature due to the energy-like-to-metal energy transfer, and blue iPr-pybox-centred photoluminescence prevailing at room temperature, as the energy back transfer to ligand is operating at higher temperatures (Figure 7b) [107]. The GdW helices with iPr-pybox, and the more expanded ind-pybox ligands, exhibit exclusively the ligand-based red phosphorescence. The NdW chains show the 4f-metal-centred NIR emission enhanced by the pybox-to-NdIII energy transfer process [108].

The rich scope of diverse lanthanides(3+) photoluminescence, and their interaction with organic 2,2’-bis(2-oxazoline) (box) organic chromophore was beautifully presented in the cyanido-bridged [LnIII(box)2(dmso)2][M(V)(CN)8]·x(solvent) (Ln = Ce–Dy, n = 2, m = 2; Ln = Ho–Yb, n = 1, m = 3; M = Mo, W) layers of a mixed 4- and 8-membered metal rings topology, showing the unusual sliding of coordination layers depending on lanthanides [109,110]. The TbW derivative, named [TbIII(box)2(dmso)2][W(V)(CN)8]·H2O (Figure 8a), reveals the excitation-dependent visible luminescence switchable between TbIII green emission induced under the deep UV excitation of the interconfigurational d-f transition of the 4f-metal centre, and red box-based phosphorescence detected for the UV excitation around 340 nm that directs the energy mainly towards the ligand excited states (Figure 8b) [109]. Similar effect was found for the analogous TbMo system while the other members of the LnMo family showed lanthanide-dependent visible and/or NIR emission. For PrMo, SmMo, EuMo and HoMo, the box-to-LnIII energy transfer induced the 4f-centred visible emission ranging from green for HoIII, orange for SmIII, to red for PrIII and EuIII. The ligand-to-metal energy transfer is also responsible for the observation of characteristic emission peaks in the NIR range for several compounds, PrMo, NdMo, SmMo, HoMo and YbMo layers [110].

6. Heteroligand Tetracyanidometallates, [MII(L)(CN)4]2− (M = Ru, Os)

Beside the homoligand polycyanidometallates, [M(CN)4]n− (x = 2, 4, 6, and 8), the photo-luminescent d-f coordination networks were prepared using the heteroligand tetracyanidometallate complexes, [MII(L)(CN)4]2− (M = Ru, Os), where two positions of the octahedron are blocked by the aromatic N,N-bidentate organic ligands, L [111–119]. Depending on the selected L ligand, exemplified by 2,2’-bipyridine (2,2’-bpy), 1,10-phenanthroline (phen), or 2,2’-bipyrimidine (bpym), these heteroligand cyanide complexes of RuII and OsII reveal the intrinsic green to red emission of the metal-to-ligand charge transfer (MLCT) origin [33]. Due to their energy position close to the red region of the visible spectrum, the MLCT excited states of [MII(L)(CN)4]2− (M = Ru, Os) ions were found to be suitable for the sensitization of NIR-emitting lanthanide(3+) ions (Figures 9 and 10, Table 5) [111–117]. It was firstly presented for the yellow emissive [RuII(2,2’-bpy)(CN)4]2− anions which combined with 4f metal ions in the aqueous solution produced the cyanido-bridged trinuclear K[LnIII(H2O)6][RuII(2,2’-bpy)(CN)4]2·9H2O (n = 7, Ln = Pr; n = 6, Ln = Er, Yb) molecules, or the two-dimensional [LnIII(H2O)4][RuII(2,2’-bpy)(CN)4]3·nH2O (Ln = Gd, Nd) layered frameworks of a mixed 4- and 12-membered metal rings topology. While the yellow MLCT emission was detected in GdRu, the NIR 4f-centred emission, enhanced by efficient RuII-to-LnIII energy transfer, was observed for PrRu, ErRu, YbRu and NdRu materials. Using the experimentally determined degree of quenching of the RuII-based emission, the rates of metal-to-metal energy transfer could be estimated. The fastest energy transfer was found for NdRu, the slowest for YbRu, while the intermediate values were assigned to PrRu and ErRu, which was rationalized on the basis of the spectral overlap between the available excited f-f states with the RuII emission band [112].
The tetracyanidoruthenate(II) complex bearing 2,2′-bipyrimidine (bpym) ligand, [Ru$^{II}$(bpym)(CN)$_4$]$^{2-}$, forms with lanthanide(3+) ions two types of coordination polymers. The lighter 4f metal ions of Nd$^{3+}$, Sm$^{3+}$ and Gd$^{3+}$ gave the helical [Ln$^{III}$(H$_2$O)$_n$(NO$_3$)$_2$][Ru$^{II}$(bpym)(CN)$_4$]$_2$ chains, while the heavier Er$^{3+}$ and Yb$^{3+}$ produced the cyanido-bridged [Ln$^{III}$$(2$H$_2$O)$_5$(NO$_3$)$_3$][Ru$^{II}$(bpym)(CN)$_4$]$_2$·5.5H$_2$O layers of a complex cross-linked square-based chains topology [113]. The red Ru$^{II}$-based emission was found for GdRu, while all the NIR-emitting Ln$^{3+}$ ions (Nd, Er, Yb) reveal their own photoluminescence, efficiently sensitized by the [Ru$^{II}$(bpym)(CN)$_4$]$^{2-}$ moieties, with the better spectroscopic overlap and resulting energy transfer efficiency for ErRu and RuYb when compared with the yellow emissive [Ru$^{II}$(2,2′-bpy)(CN)$_4$]$^{2-}$ anions. In addition, a different, expanded dinuclear bpym-bridged [(Ru$^{II}$(CN)$_4$)$_2$(bpym)]$^{4-}$ anion were prepared, and successfully applied for the construction of heterometallic d–f coordination networks. The hybrid inorganic-organic [Ln$^{III}$((H$_2$O)$_5$(NO$_3$)$_3$)$_2$][Ru$^{II}$(CN)$_4$]$_2$(bpym)]·3H$_2$O (Ln = Nd, Sm) ladder chains, and three-dimensional [Ln$^{III}$$_{1.5}$(H$_2$O)$_7$][Ru$^{II}$(CN)$_4$]$_2$(bpym)]·(NO$_3$)$_n$H$_2$O (Ln = Eu, Gd, Yb) networks of a mixed 2D cyanido-bridged, and an 1D bpym-bridged connectivity, were isolated. They revealed the analogous sensitized near-infrared 4f-metal-centred emission [113].

The bimetallic 4d–4f coordination systems involving [Ru$^{II}$L(CN)$_4$]$^{2-}$ ions offer a rich structural diversity as the blocking L ligand and four potentially bridging cyanides can result in the various low dimensional molecular systems. Moreover, the large lanthanide(3+) ions reveal high coordination numbers that the supporting organic ligands could be also introduced. It was fruitfully presented for a series of zero-, one- and two-dimensional cyanido-bridged networks based on [Ru$^{II}$(phen)(CN)$_4$]$^{2-}$ (phen = 1,10-phenanthroline) anion [114]. Using the phen molecule as the ligand coordinated both to the 4d and 4f metal centres, the cyanido-bridged [Ln$^{III}$((phen)(H$_2$O)$_3$)$_2$][Ru$^{II}$(phen)(CN)$_4$]·14H$_2$O (Ln = Nd, Gd, Er, Yb) layers of a corrugated honeycomb topology were obtained (Figure 9a). The GdRu layers reveal the strong MLCT red emission of [Ru$^{II}$(phen)(CN)$_4$]$^{2-}$ which is significantly quenched for the analogous NdRu, ErRu, and YbRu coordination frameworks due to the Ru$^{2+}$-to-Ln$^{3+}$ energy transfer process (Figure 9b). The MLCT band completely disappears for NdRu as the high density of low-lying f–f excited states of Nd$^{3+}$ accepting the energy from Ru$^{II}$. For ErRu and YbRu derivatives, the rate of d–f energy transfer, and the resulting degree of quenching of the MLCT luminescence, decreased due to the smaller number of accessible excited f–f states (Figure 9b) [114]. The similar sensitized NIR emission was found for the other phen- containing cyanido-bridged hexanuclear K$_2$[Ln$^{III}$((phen)$_2$(H$_2$O)$_2$)][Ru$^{II}$(phen)(CN)$_4$]$_4$·n(solvent) (Ln = Pr, Nd, Er, Yb) molecules, prepared under the modified synthetic conditions. The application of other ancillary oligopyridine ligands, including 2,2′:6′:2′-terpyridine (terpy) and 2,2′-bipyrimidine (bpym), resulted in the formation of one-dimensional cyanido-bridged [Ln$^{III}$(terpy)(H$_2$O)$_3$][Ru$^{II}$(phen)(CN)$_4$]$_2$·nH$_2$O (Ln = Pr, Nd, Er, Yb) and [Ln$^{III}$$_2$(bpym)(H$_2$O)$_3$][Ru$^{II}$(phen)(CN)$_4$]$_3$·n(solvent) (Ln = Nd, Er, Yb) networks of a ladder chain, and a hybrid chain of squares topologies, respectively. They all exhibit the characteristic 4f-metal-centetered NIR emission enhanced by the Ru$^{2+}$-to-Ln$^{3+}$ energy transfer. It was proved that blocking polypyridine ligands hamper the coordination of cyanides and water molecules to the 4f metal ions, not only amending the structural topology but also visibly increasing lanthanide(III)-based emission lifetimes when compared to the networks without these supporting ligands [114].
Figure 9. Crystal structure and optical properties of two-dimensional \([\text{Ln}^{III}(\text{phen})(\text{H}_2\text{O})_3]_2\)[Ru^{II}(phen)(CN)$_4$]$_4$H$_2$O (Ln = Nd, Gd, Er, Yb; phen = 1,10-phenanthroline) coordination polymers: (a) the views of the cyanido-bridged layer, and the arrangement of layers, (b) the lanthanide-dependent room temperature emission spectra in the visible range together with the related energy level diagram. Colours for the structural diagram: Ln, red; Ru, dark blue; CN$^-$, blue; phen, orange; H$_2$O, grey. Adapted from Ref. [114] with permission from The Royal Society of Chemistry.

Figure 10. Crystal structure and optical properties of three-dimensional \([\text{Ln}^{III}(\text{H}_2\text{O})_4]_2\)[[[Ru^{II}(CN)$_4$]$_3$(HAT)]$_4$H$_2$O (Ln = Nd, Gd, Yb; HAT = hexaaazatriphenylene) coordination networks: (a) the view of the whole network along the \(\hat{b}\) crystallographic direction, and the detailed view of the layered fragment of the 3D network situated within the (101) plane, (b) the lanthanide-dependent room temperature emission spectra in the visible range. Colours for the structural diagram: Ln, red; Ru, dark blue; CN$^-$, blue; HAT, orange; H$_2$O, grey [116]. Reprinted with permission from J. Am. Chem. Soc. 2007, 129, 11491–11504. Copyright 2007 American Chemical Society.
### Table 5. Photoluminescent coordination compounds containing lanthanide ions and heteroligand tetracyanidometallates.

| Compound | Structural type | Luminescent property | Ref. |
|----------|-----------------|----------------------|------|
| [K(LnIII(H₂O)₆)[Ru(II)(2,2'-bipy)(CN)₄]·9H₂O] (n = 7, Ln = Pr; n = 6, Ln = Er, Yb) | 0D trimolecular, 2D mixed 4- and 12-membered metal rings | NIR LnⅢ (Ln = Pr, Er, Yb) emission with RuII-to-LnⅢ ET.² | [111,112] |
| [LnIII(H₂O)₆][Ru(II)(2,2'-bipy)(CN)₄]·12H₂O (Ln = Gd, Nd) | | NIR NdⅢ emission (NdRu) with RuⅡ-to-NdⅢ ET. Yellow | [112] |
| [LnIII(H₂O)₅NO₃][Ru(II)(4bpy)(CN)₄]·4MeOH (Ln = Nd, Sm, Gd) | 1D helical chain | NIR NdⅢ emission (NdRu) with RuⅡ-to-NdⅢ ET. Red | [113] |
| | | [Ru(4bpy)(CN)₄]²⁺ emission in GdRu (MLCT).³ | |
| [LnIII(H₂O)₅(NO₃)₂][Ru(II)(4bpy)(CN)₄]·4MeOH (Ln = Nd, Sm, Gd) | 2D cross-linked | NIR LnⅢ (Ln = Er, Yb) emission with RuⅡ-to-LnⅢ ET. Yellow | [113] |
| [LnIII(H₂O)₅(NO₃)₃][Ru(II)(CN)₄]·3(H₂O)·4MeOH (Ln = Nd, Sm) | 1D hybrid ladder chain | NIR NdⅢ emission in NdRu with RuⅡ-to-NdⅢ ET. | [113] |
| [LnIII(H₂O)₅(CN)₄][Ru(II)(CN)₄]₂(Bpy)(NO₃)·nH₂O (Ln = Eu, Gd, Yb) | 3D hybrid P₂O₅ network | NIR YbⅢ emission (YbRu) with RuⅡ-to-YbⅢ ET. Red [Ru(bpy)(CN)₄]²⁺ emission in GdRu (MLCT). | [113] |
| K₂[LnIII(phen)₂(H₂O)₂][Ru(II)(phen)](CN)₄·n(solvent) (Ln = Pr, Nd, Er, Yb) | 0D hexanuclear | NIR LnⅢ (Ln = Nd, Er, Yb) emission with RuⅡ-to-LnⅢ ET. | [114] |
| [LnIII(terpy)(H₂O)₅][Ru(II)(phen)](CN)₄·n(solvent) (Ln = Pr, Nd, Er, Yb) | 1D ladder chain | NIR LnⅢ (Ln = Nd, Er, Yb) emission with RuⅡ-to-LnⅢ ET. | [114] |
| [NdIII(H₂O)₅][Ru(II)(CN)₄]·3H₂O | 1D hybrid chain of squares | NIR YbⅢ emission with RuⅡ-to-YbⅢ ET. | [114] |
| [LnIII(H₂O)₅][Ru(II)(HAT)(CN)₄]·nH₂O (Ln = Nd, Sm, Eu, Gd) | 2D hybrid of metal rings | NIR NdⅢ emission (NdRu) with RuⅡ-to-NdⅢ ET. Red [Ru(phen)(CN)₄]²⁺ emission in GdRu (MLCT). | [115] |
| | | | |
| [YbIII(H₂O)₅(NO₃)₃][Ru(II)(HAT)](CN)₄·6H₂O | 1D ladder chain | NIR YbⅢ emission with RuⅡ-to-YbⅢ ET. | [116] |
| [YbIII(H₂O)₅(NO₃)₃][Ru(II)(HAT)](CN)₄·2H₂O | 2D hybrid 6-membered metal rings | NIR YbⅢ emission with RuⅡ-to-YbⅢ ET. | [116] |
| [LnIII(H₂O)₅][Ru(II)(CN)₄]·12H₂O (Ln = Nd, Gd, Tb) | 3D pillared network | NIR LnⅢ (Ln = Nd, Gd, Tb) emission with RuⅡ-to-LnⅢ ET. | [116] |
| [GaIII(H₂O)₆][Os(III)(MeOH)](O₂)(phen)·4H₂O (CN)₄·1.5H₂O | 2D 12-membered metal rings | Red [Os(phen)(CN)₄]²⁺ emission (MLCT). | [117] |
| Na₂[LnIII(phen)₂(H₂O)₆][NO(phen)](CN)₄·4MeOH·17H₂O (Ln = Pr, Nd, Er, Yb) | 0D hexanuclear | NIR LnⅢ (Ln = Nd, Er, Yb) emission with OsⅢ-to-LnⅢ ET. | [117] |
| [LnIII(phen)₂(H₂O)₆][NO(phen)](CN)₄·4MeOH·17H₂O (Ln = Pr, Nd, Er, Yb) | 1D hybrid chain of squares | NIR LnⅢ (Ln = Nd, Er, Yb) emission with OsⅢ-to-LnⅢ ET. | [117] |
| [LnIII(phen)₂(H₂O)₆][NO(phen)](CN)₄·4MeOH·17H₂O (Ln = Pr, Nd, Er, Yb) | 2D mixed 3-, 4-, 6-, and 14-membered metal rings | NIR NdⅢ emission with OsⅢ-to-NdⅢ ET. Red | [117] |
| | | | |
| K₂[LnIII(H₂O)₅][Ru(II)(bpy)(CN)₄]·3H₂O·8MeOH (Ln = Pr, Nd, Sm, Eu) | 1D chain of squares | Orange [Ru(bpy)₂(bpy)(CN)₄]²⁺ emission (MLCT); selective sensitization by gaseous amine molecules. | [118,119] |

¹ 2,2'-bipy = 2,2'-bipyridine; ² ET = energy transfer; ³ MLCT = metal-to-ligand charge transfer; ⁴ HAT = hexaalatrinaphenylene; ⁵ bpy = 2,2'-bipyridimidine; ⁶ phen = 1,10-phenanthroline; ⁷ terpy = 2,2'-6',2''-terpyridine; ⁸ ‘Bubpy = 4,4''-di-tert-butyl-2,2'-bipyridine.

The particularly efficient sensitization of both Nd³⁺ and Yb³⁺ was achieved for the heterometallic d-f coordination networks based on tetracyanidometallates(II) bearing the expanded aromatic system of hexaalatrinaphenylene (HAT) (Figure 10) [115,116]. The simplest metal-cyanide complex of this family, [Ru(II)(CN)₄](HAT)²⁺ was combined with lanthanide(III) ions giving the hybrid layered [LnIII(H₂O)₅][Ru(II)(HAT)(CN)₄]·nH₂O framework of a 12-membered metal rings topology for larger lanthanides of Nd, Sm, Eu, Gd, or the one-dimensional cyanido-bridged [YbIII(H₂O)₅(NO₃)₃][Ru(II)(HAT)(CN)₄]·2(NO₃)·6.5H₂O ladders bearing smaller Yb³⁺ ions. Using the other cyanide precursor of [Ru(II)(CN)₄](HAT)⁴⁻ with two tetracyanidometallates(II) attached to a single HAT, the very different hybrid [YbIII(H₂O)₅(NO₃)₃][Ru(II)(CN)₄](HAT)²⁺·20H₂O layers of 6-membered metal rings topology...
were synthesized. The most expanded [(RuII(CN)4)3(HAT)]3— anion, offering three RuII centres linked by a single HAT, and 12 potentially bridging cyanide ligands, was mixed with lanthanide(3+) ions producing the three-dimensional [LnIII(H2O)12][(RuII(CN)4)3(HAT)]13H2O (Ln = Nd, Gd, Yb) pillared network. They are constructed of the hybrid HAT-cyanide heterometal layers connected in the perpendicular direction by additional cyanide linkages (Figure 10a) [115,116]. When Gd3+ ion was used as the lanthanide centre, all the RuII(HAT)-containing networks reveal the strong MLCT photoluminescence in the red range, which was substantially quenched by the insertion of NIR-emitting Nd3+ and Yb3+ ions. As a result, the strong near-infrared emission was observed, while the residual RuII-based emission was around 100 times weaker than for the GdRu compounds (Figure 10b) [116].

The heteroligand [OsII(phen)(CN)4]2— ion, built of 5d OsII metal centre, reveals the red MLCT emission, shifted towards lower energy as compared to [RuII(phen)(CN)4]2—, which was presented for the layered [GdIII(H2O)4(MeOH)][OsII(phen)(CN)4]1.5·4H2O framework of a 12-membered metal rings topology [117]. This makes the tetracyanidoosmate(II) an even more efficient sensitizer for NIR-emitting lanthanide(3+) ions than the related tetracyanidoruthenate(II) due to the expected better spectroscopic overlap of Os2+­-based emission with the absorption peaks of the 4f metal ions. This prediction was checked for the family of cyanido-bridged coordination systems, including hexanuclear Na2[LnIII(phen)2(H2O)2]2[OsII(phen)(CN)4]12·4MeOH·17H2O (Ln = Pr, Nd, Er, Yb) clusters, one-dimensional [LnII2(bpym)(H2O)]2[OsII(phen)(CN)4]12·MeOH·13H2O (Ln = Er, Yb) hybrid chain of squares, and the intricate [NdII4(bpym)2(H2O)12(MeOH)][OsII(phen)(CN)4]4·6MeOH·19.5H2O coordination layers. For all these species, the NIR lanthanide-based emission was sensitized by the Os2+­-to-Ln3+ energy transfer of distinguishable rates depending on 4f metal ion. The rates of Os2+­-to-Ln3+ energy transfer were found to be an order of magnitude faster than the rates of RuII­-to-Ln3+ energy transfer previously observed in similar heterometallic d-f cyanido-bridged networks [117].

The innovative luminescent functionality of heteroligand tetracyanidometallates was shown for the [RuII(Bubpy)(CN)4]2— ion bearing the tert-butyl derivative of 2,2’-bipyridine [118,119]. This anion embedded in the bimetallic K[LnIII(H2O)4][RuII(Bubpy)(CN)4]2·8H2O (Ln = Pr, Nd, Sm, Eu) chain of [Ln2Ru2] molecular squares exhibits the orange emission of a typical metal-to-ligand charge transfer (MLCT) origin. The intensity of this photoluminescence detected for the bulk solid sample, and the polymeric film, was reported to be strongly dependent on the presence of gaseous amine molecules, which additionally sensitize the RuII-based emission. Therefore, this material found the analytical application as an efficient and sensitive chemodosimetric detector of biogenic amine odorants including histamine, putrescine, spermidine, and ammonia, indicating a great potential of the polycyanidometallate-based d-f coordination networks in chemical and biochemical sensing based on the spectrofluorometric detection [119].

7. Other Cyanide-Containing Building Blocks

The photoluminescent cyanido-bridged coordination networks incorporating trivalent lanthanide ions were achieved by a few non-typical cyanide-building blocks (Figures 11 and 12, Table 6) [120–123]. For instance, the solvolothermal reaction of a chemically modified salen-type ligand, N,N’-ethylene bis[4-(ethylamino)salicylideneimine] (L1), with erbium(III) chloride and copper(I) cyanide in a non-aqueous methanol-acetonitrile solution resulted in the isolation of an unprecedented [ErIII2CuII2(L1)2(Cl)2][CuI4(CN)6(MeCN)4] supramolecular network. It is composed of the two-dimensional homometallic CuI-based cyanido-bridged layers of a 10-membered metal rings topology with the trinuclear salen-ligand-bridged [ErIII2CuII2L12Cl2] clusters occupying the interlayer space (Figure 11a) [120]. This material reveals NIR Er3+-centred emission under the visible light excitation, slightly weakened when compared with the reference cyanide-free [ErIII2(H2L)4] molecule due to the partial Er3+-to-CuI f-d energy transfer (Figure 11b). This work showed that a cuprous cyanide network serves as a stabilizing agent for the formation of unique NIR-emitting [ErIII2CuII2L12Cl2] clusters, non-isolable under cyanide-free synthetic conditions [120].
The rare organometallic heteroligand dicyanidoiridate(III) anion, $[\text{Ir}^{III}(ppy)_2(CN)_2]^{-}$, with four coordination sites blocked by two anions of 2-phenylpyridine (ppy), was also tested as a novel cyanide-containing metal complex for d–f photoluminescent materials [121]. The self-assembly of this specific dicyanido metallate with the excess of lanthanide(3+) ions resulted in a series of cyanido-bridged tetranuclear $[\text{Ln}^{III}(H_2O)_2(\text{MeCN})(\text{NO}_3)_2]_2[\text{Ir}^{III}(ppy)_2(\text{CN})_2]_2\cdot 4\text{MeCN}$ ($\text{Ln} = \text{La, Pr, Nd}$), $[\text{N}(\text{PPh}_3)_2]_2[\text{Eu}^{III}_2(H_2O)(\text{NO}_3)_6][\text{Ir}^{III}(ppy)_2(\text{CN})_2]_2\cdot 5\text{MeCN}$, and $[\text{Gd}^{III}(\text{NO}_3)_2(H_2O)_2]_2[\text{Ir}^{III}(ppy)_2(\text{CN})_2]_2$ molecules which structural details depend on the attached lanthanide(3+) ions. In all these materials, the broad green $[\text{Ir}^{III}(ppy)_2(\text{CN})_2]^{-}$-based emission was detected, the strongest for the UV-emissive Gd$^{3+}$, and much weaker for visible-light-emissive Eu$^{3+}$, and NIR-emitting Nd$^{3+}$. It suggested the presence of Ir$^{3+}$-to-Eu$^{3+}$ and Ir$^{3+}$-to-Nd$^{3+}$ energy transfer processes, but the sensitized Eu$^{3+}$ or Nd$^{3+}$ could not be detected indicating the additional quenching effect, presumably involving the water molecules existing in the crystal structure [121].

**Figure 11.** Crystal structure and optical properties of supramolecular $[\text{Eu}^{III}_2\text{Cu}^{II}_2(L_1)_2(\text{Cl})_2]$ $[\text{Cu}_2^4(\text{CN})_5(\text{MeCN})_4]_2$ ($L_1 = \text{N,N’-ethylene bis[4-(ethylamino)salicylideneimine anion}$) network: (a) the structural views of two main components, including the fragment of the cyanido-bridged CuI-based layer and the $[\text{ErIIICuI}_2]$ molecule, and their arrangement in the supramolecular network, (b) the room temperature excitation and emission spectra of the reference cyanide-free $[\text{ErIIICuII}_2(L_2)_2]$ molecule (black and purple lines), and the presented supramolecular network (red and light purple lines). Colours for the structural diagram: Er, red; Cu, dark blue; CN$^{-}$, blue; L1, orange; Cl$^-$, light green. Adapted from Ref. [120] with permission from The Royal Society of Chemistry.
excitation pathways \[121\]. Thus, \([\text{B}^{\text{III}}(\text{CN})_4]^-\) serves only as a supporting inorganic ligand without the significant emission or sensitizing effect. Much more pronounced role in the photoluminescent effect is played by its structurally modified derivative, that is pentafluoroethyltricyanidoborate(III) ion, \([\text{C}_2\text{F}_5\text{B}^{\text{III}}(\text{CN})_3]^-\)\[123\]. This anionic building block combined with lanthanide(3+) ions resulted in the formation of a hydrated cyanido-bridged \([\text{Ln}^{\text{III}}{\text{C}_2\text{F}_5\text{B}(\text{CN})_3}^3(\text{H}_2\text{O})_3]\) (\(\text{Ln} = \text{La}, \text{Eu}, \text{Ho}\)) layered framework showing the topology of cross-linked \({\text{Ln}}_2\text{B}_2\)-square-based chains (Figure 12a). Upon thermal dehydration, this material could be transformed into the anhydrous three-dimensional cyanido-bridged \([\text{Ln}^{\text{III}}{\text{C}_2\text{F}_5\text{B}(\text{CN})_3}] (\text{Ln} = \text{La}, \text{Eu}, \text{Ho})\) hexagonal network built of nine-coordinated

![Crystal structure and optical properties of two-dimensional \([\text{Ln}^{\text{III}}(\text{C}_2\text{F}_5\text{B}(\text{CN})_3)_3(\text{H}_2\text{O})_3]\) (\(\text{Ln} = \text{La}, \text{Eu}, \text{Ho}\)) coordination polymer (a,c), and the related anhydrous three-dimensional \([\text{Ln}^{\text{III}}(\text{C}_2\text{F}_5\text{B}(\text{CN})_3)_3]\) (\(\text{Ln} = \text{La}, \text{Eu}, \text{Ho}\)) coordination network (b,c): (a) the views of the single layer, and the arrangement of layers, (b) the views of the 3D cyanido-bridged network along \(c\) and \(b\) crystallographic axes, (c) the room temperature excitation and emission spectra of the 2D \(\text{LnB}\) layer (solid lines), and 3D \(\text{LnB}\) network (dotted lines). Colours for the structural diagram: \(\text{Ln}, \text{red}; \text{B}, \text{dark blue}; \text{CN}^-, \text{blue}; \text{C}_2\text{F}_5^-, \text{orange}; \text{H}_2\text{O}, \text{grey}\)\[123\]. Reprinted with permission from Inorg. Chem. 2017, 56, 2278–2286. Copyright 2017 American Chemical Society.

Non-heterometallic, however, cyanido-bridged photoluminescent coordination networks were obtained by combining trivalent lanthanide ions with inorganic tetracyanidoborate, \([\text{B}^{\text{III}}(\text{CN})_4]^-\) anionic building block, and its heteroligand pentafluoroethyltricyanidoborate(III), \([\text{C}_2\text{F}_5\text{B}^{\text{III}}(\text{CN})_3]^-\) derivative \[122,123\]. The rare-earth metal oxide or hydroxides mixed with tetracyanidoboronic acid, \(\text{H}[\text{B}(\text{CN})_4]\cdot n\text{H}_2\text{O}\) produced the cyanido-bridged dinuclear \([\text{Ln}^{\text{III}}(\text{H}_2\text{O})_7][\text{B}(\text{CN})_4]2[\text{B}(\text{CN})_4]\) (\(\text{Ln} = \text{Tb}, \text{Dy}\)) molecules where tetracyanidoborate(III) serves as a ligand, and a crystallization counterion. The TbB and DyB compounds reveal the intense sharp peaks of their characteristic visible photoluminescence, realized by direct intrashell \(f\)-\(f\) excitation, and interconfigurational \(f\)-\(d\) excitation pathways \[121\]. Thus, \([\text{B}^{\text{III}}(\text{CN})_4]^-\) serves only as a supporting inorganic ligand without the significant emission or sensitizing effect. Much more pronounced role in the photoluminescent
effect is played by its structurally modified derivative, that is pentafluoroethyltricyanidoborate(III) ion, [C₂F₅BF⁺(CN)₃]⁻ [123]. This anionic building block combined with lanthanide(3+) ions resulted in the formation of a hydrated cyanido-bridged [LnIII(C₂F₅BF(CN)₃)(H₂O)₃](Ln = La, Eu, Ho) layered framework showing the topology of cross-linked [Ln₂B₂]-square-based chains (Figure 12a). Upon thermal dehydration, this material could be transformed into the anhydrous three-dimensional cyanido-bridged [LnIII(C₂F₅BF(CN)₃)](Ln = La, Eu, Ho) hexagonal network built of nine-coordinated Ln³⁺ ions which coordination sphere is fully occupied by cyanide ligands of bridging [C₂F₅BF⁺(CN)₃]⁻ anions (Figure 12b). The LaB anhydrous network exhibits the broad greenish-blue UV-light-induced photoluminescence of tricyanidoborate(III) anions. This emission is also observed for HoB analogues but the intense reabsorption effect was detected as the several characteristic f-f lines situated within the broad emission band of the anion. It suggested the lack of efficient [C₂F₅BF⁺(CN)₃]⁻ to-Ho³⁺ energy transfer. On the contrary, the sensitization effect was found for the hydrated and anhydrous EuB networks that show the strong 4f-metal-centred emission, and the complete quenching to-Ho³⁺ emission under direct f-f, and f-d excitations. [121]

Table 6. Photoluminescent coordination compounds composed of lanthanide ions and other cyanide-containing building blocks.

| Compound | Structural type | Luminescent property | Ref. |
|----------|-----------------|----------------------|------|
| [ErIII(CuII)(L1)(Cl)]²⁺| 2D Cu-cyanide | NIR Eu³⁺ emission under the visible excitation. | [120] |
| [CuIII(CN)₃(MeCN)₄] | 10-membered metal rings, and trinuclear [ErCu]³⁺ counterions | | |
| [LnIII(H₂O)₂(MeCN)(NO₃)₂]²⁺, [IrIII(ppy)₂(CN)₃] | 4MeCN (Ln = La, Pr, Nd) | 0D tetranuclear molecule | Green [IrIII(ppy)₂(CN)₃]⁻ emission in NdIr, weakened by partial Ir³⁺-to-Nd³⁺ ET.⁵ | [121] |
| [N(PPPh₃)₂][II(ppy)²(CN)₂] | 5MeCN | 0D tetranuclear molecule | Green [IrIII(ppy)₂(CN)₃]⁻ emission, weakened by partial Ir³⁺-to-Eu³⁺ ET.⁵ | [121] |
| [LnIII(ppy)₂(CN)₃] | 2D tetranuclear molecule | Green [IrIII(ppy)₂(CN)₃]⁻ emission. | [121] |
| [LnIII(H₂O)₇][B(CN)₄]²⁺ | 2D cross-linked square-based chains | Green Eu³⁺ emission in EuB with [C₂F₅BF(CN)₃]⁻ energy transfer. | [123] |
| [LnIII(C₂F₅BF(CN)₃)] | square-based chains | Red Eu³⁺ emission in EuB with | [123] |
| [LnIII(C₂F₅BF(CN)₃)] | 3D network | Greenish-blue [C₂F₅BF(CN)₃]⁻ emission in LaB and HoB with reabsorption effect in HoB. Red Eu³⁺ emission in EuB EuB with | [123] |

1 L1 = N,N’-ethylene bis[4-(ethylamino)salicylideneimine] anion; ² ppy = anion of 2-phenylpyridine; ³ ET = energy transfer.

8. Conclusions

We have presented a detailed overview of lanthanide photoluminescence in coordination networks based on polycyanidotellurates. To best of our knowledge, this is the first systematic catalogue of the state of the art in the investigation of photoluminescent d–f cyanido-bridged systems.

Selected groups of polycyanidotellurates are suitable for construction of photoluminescent coordination frameworks that can explore intrinsic optical properties of lanthanide(3+) ions. Among them, the dicyanidotellurates of Au⁴⁺ and Ag⁴⁺ are luminescent in the visible range due to the metal-to-ligand charge transfer (MLCT) transitions. Such emission, found in 4d/5d–4f coordination polymers, and facilitated by the presence of Au–Au interactions, was shown to be dependent on pressure and temperature. Dicyanidotellurates are good sensitizers for visible emission of 4f-metal ions, and the resulting emission can be tuned by external stimuli such as pressure, temperature, or light excitation. Thus, the multi-coloured tunable photoluminescence is the main
application of [M(\text{CN})_2]^- based d-f systems, taking also into account the possible insertion of organic ligands, introducing additional energy transfer pathways, or separate emission components. Similar luminescence was achievable for tetracyanidoplatinate(II) ions revealing the visible emission due to the metal-to-metal-to-ligand charge transfer (MMLCT) transitions related to the Pt-Pt interactions. This emission is dependent on temperature, pressure, and light polarization due to the anisotropic character of these square planar polycyanido- metallas. [Pt(\text{CN})_4]^{2-} is a good sensitizer for some lanthanide(3+) ions, including Eu^{3+}, Sm^{3+}, or Tb^{3+}. The tetracyanidoplatinate(II)-based systems offer a structural diversity due to the ancillary organic ligands, modifying the 4f-metal-centred emission.

The octahedral hexacyanidometallates of Co^{III} and Cr^{III} are luminescent in the red and NIR ranges, respectively. They can enhance NIR lanthanide(3+) emission by the energy transfer. [Co(\text{CN})_6]^{3-} can also sensitize visible 4f-metal-centred luminescence. They offer six cyanides to create extended networks, and leave the coordination sites on lanthanides that can be occupied by organic ligands. Thus, various optical functionalities were achieved, as exemplified by white light emission of Dy^{III}−Co^{III} compound, excitation- and composition-switchable multi-coloured emission in Eu^{III}−Tb^{III}−Co^{III} chains, and dual-donor NIR emission in Nd^{III}−Cr^{III} and Yb^{III}−Co^{III} networks.

Despite their non-luminescent character, octacyanidometallates of Mo^{IV/V} and W^{IV/V} were fruitfully applied in the synthesis of photoluminescent d-f coordination systems. Due to the optical transparency in the Vis–NIR range, their cyanido-bridged networks with lanthanides could reveal the rich spectrum of intrinsic properties of 4f metal ions, and the sensitization effects on supporting organic ligands. [M(\text{CN})_6]^{3-} ions offer a great structural flexibility, so a number of d-f coordination networks could be rationally designed. As a result, such attractive functionalities as thermal switching of blue–red emission in Eu^{III}−W^{V} helices, excitation-tunable green–red luminescence in Tb^{III}−W^{V} layers, or humidity sensor based on red emissive Eu^{III}−Mo^{IV} network, were reported.

The heteroligand tetracyanidometallates of Ru^{II} and Os^{II}, [M(\text{CN})_4(L)]^{2-} are photoluminescent in the visible range due to the metal-to-ligand charge transfer (MLCT) transition. These anions are good sensitizers for NIR-emitting lanthanide(3+) ions. Their emission energy can be tuned by the blocking ligand L, thus, the selection of appropriate organic part governs the efficiency of the Ru^{2+}/Os^{2+}−to-Ln^{3+} energy transfer process. Metal center also plays a vital role, and the Os^{II}-based cyanide complexes were proved to be better sensitizers for NIR emission than the analogous Ru^{II} species. The ancillary organic ligands can be, here, inserted to amend the structural features, and facilitate the characteristics of energy transfer. Lately, the heteroligand tetracyanidoruthenate(II) embedded in d-f cyanido-bridged networks has successfully been applied in the chemodosimetric detection of biogenic amines utilizing a selective sensitization of its visible MLCT emission.

Among other cyanide-containing building blocks, the cuprous cyanide networks was found to stabilize NIR-emitting salen-based Er^{III}−Cu^{III} clusters, while the green emissive heteroligand dicyanidoiridate(III) bearing two blocking bidentate ligands were tested as possible sensitizers for lanthanide(3+) luminescence. In addition, the inorganic tetracyanidoborate anion was used for the construction of photoluminescent Tb^{III} and Dy^{III}-based molecules, however, without a strong impact on the emission properties. On the contrary, its modified analogue, pentafluoroethyli-tricyanidoborate(III) ion, [C_2F_5B(\text{CN})_3]^- is greenish-blue emissive, and efficiently sensitizes the Eu^{3+} luminescence within the cyanido-bridged two- or three-dimensional coordination networks.

In summary, we have gathered all reported photoluminescent polycyanidometallate-based coordination systems incorporating lanthanide ions. We have shown that cyanide metal complexes are useful for the preparation of emissive solids revealing such functionalities as lanthanide- centred Vis–NIR photoluminescence, sensitized through metal-to-metal and/or ligand-to-metal energy transfer effects, and multi-coloured emission tuned by external stimuli of excitation wavelength, temperature, and pressure. There is still several challenges in this area, including the detailed investigation of structural and electronic features governing the sensitization process involving cyanide complexes, and the preparation of novel luminescent polycyanidometallates of improved optical properties, and affinity to lanthanide(3+) ions. The particularly promising future work comprises of searching for
multifunctional photoluminescent materials that can exhibit the undiscovered physical cross-effects, as polycyanidometallate-based heterometallic coordination networks showed a diversity of physical properties, such as magnetic anisotropy leading to slow magnetic relaxation, magnetic ordering, spin transitions, catalytic activity, ferroelectricity, and ionic conductivity, that in some cases lead to the unprecedented physical phenomena [38–42,50].

Acknowledgments: This work was financed by the National Science Centre, Poland within the SONATA 11 project, grant no. 2016/21/D/ST5/01634. B.S. is grateful for the financial support of the National Science Centre, Poland within the OPUS-8 project, grant no. 2014/15/ST5/04465.

Author Contributions: S.C.: conception, literature search, and manuscript preparation; M.W.: literature structural search and manuscript preparation; B.S.: review of the manuscript and contribution to the manuscript preparation.

Conflicts of Interest: The authors declare no conflict of interest.

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