# Combining electron transfer, spin crossover, and redox properties in metal-organic frameworks

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Hofmann coordination polymers (CPs) that couple the well-studied spin transition of the Fe<sup>II</sup> central ion with electron-responsive ligands provide an innovative strategy toward multifunctional metal-organic frameworks (MOFs). Here, we developed a 2D planar network consisting of metal-cyanide-metal sheets in an unusual coordination mode, brought about by infinitely π-stacked redox-active bipyridinium derivatives as axial ligands. The obtained family of materials show vivid thermochromism attributed to electron transfer and/or electronic spin state change processes that can occur either independently or concomitantly. Importantly, the redox activity of the ligands within the structure leads to the quasi-reversible electrochemical reduction reaction on a spin-crossover complex at solid state. These observations have been confirmed via temperature-dependent single-crystal X-ray diffraction, magnetic measurements, Mössbauer, EPR, optical and vibrational spectroscopies as well as quantum chemical calculations.

Multi-switchable and readable molecules able to reversibly tune their intrinsic properties in response to one or several external stimuli have received much attention as potential components in a great variety of applications, e.g. memory devices<sup>1</sup>, sensors<sup>2,3</sup>, or catalysts<sup>4</sup>. In general, multifunctional compounds consist of several bistable building blocks, each able to respond to external stimuli (such as light, temperature, electrochemical or magnetic input) $5,6$ . Among the different building blocks, bipyridinium units<sup> $7-11$  $7-11$  $7-11$ </sup> are especially promising because of their excellent electrochemical properties due to their electron-deficient character: the one-electron reduction in diluted solutions yields the paramagnetic bipyridinium radical cation, whereas it induces the dimerization of the molecules in a concentrated solution. In order to take advantage of the redox properties, bipyridinium derivatives have been successfully included in a large variety of systems spanning the fields of supramolecular chemistry, coordination chemistry, hostguest compounds, electrochromic materials etc. aiming to induce an inter or intra-molecular electron transfer (ET) by applying an external stimulus (temperature, light irradiation or pressure) $12-18$ . As an example, Coe et al.<sup>[19](#page-10-0)</sup> reported the possibility for pentacyano-ironpyridinium derivatives to undergo an ET due to the electron donor and acceptor character of the building blocks. Much more common are examples of coordination polymers where the observed piezo, photo, or thermal ET is due to the presence of an electron-deficient viologen derivative, and a carboxylic acid acting as the electron-rich counterpart $17,20,21$ . In such configurations, the ET occurs through hydrogen bonding, the most likely electron donor being the uncoordinated carboxylate oxygen atom. In general, the key structural factors favoring ET pathways are: (i) short distances between donor and acceptor, where the ET transfer will occur through either space or hydrogen bonds and (ii) planar structures to allow for electron delocalization. However, one should note that the structural changes before and after the ET observed in the literature are subtle.

Metal organic frameworks (MOFs) on a bipyridinium basis can be excellent candidates to observe an ET due to their structural arrangement. Recently, Guo et al.<sup>12</sup> reported a 2D semiconductor MOF composed by π-stacked redox active N-methylpyridinium cations,

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sandwiched by cyanide-bridged layers that shows light and temperature-induced color change with the formation of stable radicals even in an ambient atmosphere. In this case, the observed thermoand photo-activated ET within these materials involves the pyridinium unit as acceptor and the uncoordinated CN- moiety as donor. These MOF structures are closely related to Prussian blue and/or Hofmann clathrate derivatives which offer excellent structural platforms to establish multifunctionality, due to their intrinsic magnetic properties: photomagnetism, magnetic ordering, valence tautomerism and/or a spin crossover (SCO)<sup>22-[26](#page-11-0)</sup>.

In general, Hofmann clathrates are formed by three major building blocks, first off, a transition metal center, second, a ligand (L) and lastly, a cyanometallate anion  $[M(CN)_x]^n$ <sup>,</sup> (x = 2 or 4, n = 1 or 2). Usually, this combination affords 2D planar metal−cyanide−metal sheets completed by a mono- or bidentate neutral organic ligand L, most often ammonia<sup>27</sup>, pyridine<sup>28-32</sup>, triazole<sup>33-[37](#page-11-0)</sup> or pyrazine<sup>[38](#page-11-0)-[40](#page-11-0)</sup> derivatives, with a general formula of  ${Fe(L)_x[\mu_4 \cdot M'(CN)_4]}$  (x = 1 or 2; M'= Ni, Pd or Pt) or  ${Fe(L)_x[\mu_2\text{-}M''(CN)_2]}$  (x = 1 or 2; M''= Cu, Ag or Au)<sup>28</sup>. The modulation of the ligand field around the iron center through the different units leads to the introduction of the SCO properties. To our knowledge, at least three exceptions to this general formula have been described in literature:  $i$ ) if the ligand is a strong chelate, for example a quinoline derivative (aqin), 1D chains of  $[Fe(aqin)<sub>2</sub>(\mu<sub>2</sub>-M(CN)<sub>4</sub>)]$  (M = Ni<sup>II</sup> and Pt<sup>II</sup>) are obtained, where [M(CN)<sub>4</sub>]<sup>2</sup> acts as a bridge between the iron sites<sup>41</sup>; *ii*) if the ligand is tetradentate, a  $\{[Fe(\mu_4-bztpy)\mu_2 -$ Pt(CN)4]·0.5bztpy} structure emerges, where 2D sheets, formed by the ligand and the Fe<sup>II</sup> metal center, are again interconnected by  ${\rm [M(CN)_4]}^2$ building blocks<sup>42</sup>, and *iii*) very recently, Real et al.<sup>43</sup> described two isomorphous structures, where the ligand carries a positive charge due to a spontaneous protonation during the crystallization process. The authors suggested that the positive charge could be responsible for an additional class of Hofmann-clathrates with the general formula  ${Fe(L)}_2[u_2-M(CN)_4]$ . It should be noted that in all of the above cases, the specific properties (electronic, steric, etc.) of the ligands are responsible for the final structure of these modified Hofmann-type clathrates, which, nevertheless, maintain a SCO behavior. Thus, we think that Hofmann-type clathrates offer an excellent structural platform to establish multifunctionality through introduction of a redoxactive ligand, such as a bipyridinium-type derivative.

In this work, we explore this idea by synthesizing a large family of non-innocent ligands, which contain three functional building blocks (Fig. 1a; see Supplementary Figs 1–5 for characterization), the most important being a monocationic pyridinium unit that is responsible for the redox activity<sup>12,[44](#page-11-0)-[47](#page-11-0)</sup>. Attached to it, an aryl group allows for the finetuning of the delocalized electron density of the ligand by modifying the substituent R in para position. Lastly, a pyridine group is added to

ensure the coordination to the iron center. The pyridinium moiety contains an electropositive, quaternary nitrogen atom that can be reduced to the highly air-sensitive neutral (radical) species (Fig. 1b). As demonstrated through several examples in the literature, the stability of the radical state can be largely improved by e.g. incorporating it in a MOF structure. In the case of the compound reported by Guo et al. $12$ the stabilization of the radical is due to π−π and cation-π interactions within the network. With this idea in mind, the abovementioned redoxactive ligands were used to synthesize a family of Hofmann-type MOFs, which, as we show below, host a wealth of electronic phenomena, including both SCO and ET processes as well as redox activity.

### Results

In the present work, we report the synthesis and characterization of the first redox active spin crossover coordination polymers belonging to this family. These compounds are isostructural and consist of 2D extended sheets with the molecular formula  $\{Fe(\mathbf{R}\cdot\mathbf{pbpy}^*)\}$ [µ2- $M(CN)<sub>4</sub>$ ]<sub>2</sub>·2H<sub>2</sub>O} (R being methyl (1, CH<sub>3</sub>), bromo (2, Br), ester (3, COOCH<sub>3</sub>) or nitro (4, NO<sub>2</sub>),  $\mathbf{R}$ -pbpy<sup>+</sup>= 1-(4-R-phenyl)–4,4'-bipyridinium and M<sub>1</sub> = Ni<sup>2+</sup> (**a**) or M<sub>2</sub> = Pt<sup>2+</sup> (**b**)). The {Fe(**R-pbpy**<sup>+</sup>)<sub>2</sub>[ $\mu$ <sub>2</sub>-M(CN)<sub>4</sub>]<sub>2</sub>·2H<sub>2</sub>O} Hofmann CPs were synthesized as microcrystalline powders and as single crystals. The direct powder precipitation of the compounds is obtained when  $K_2[M(CN)_4] \times H_2O (M = Ni^{2+}$  or  $Pt^{2+}$ ) is added to an EtOH/ H<sub>2</sub>O (1:1) solution containing the ligand and  $Fe(BF<sub>4</sub>)<sub>2</sub>·6H<sub>2</sub>O$ . The purity of the products has been assessed by elemental analysis (Supplementary Methods) and powder X-ray diffraction (XRD) (Supplementary Figs. 9–16). Single-crystals suitable for XRD analysis were obtained by slow crystallization in an H-tube from the diffuse reaction of  $Fe(BF<sub>4</sub>)<sub>2</sub>·6H<sub>2</sub>O$ , the ligand  $R\cdot pbpy^+$  (R=CH<sub>3</sub>, Br or COOCH<sub>3</sub>) and  $K_2[Pt(CN)_4]$  in a molar ratio of 1:2:2 (see Supplementary Tables 9, 10 & 12 for compounds **1b, 2b & 3b** respectively).

The unit cell of  ${Fe}$ (CH<sub>3</sub>-pbpy<sup>+</sup>)<sub>2</sub>[µ<sub>2</sub>-Pt(CN)<sub>4</sub>]<sub>2</sub>·2H<sub>2</sub>O} (1b) (P2<sub>1</sub>/c monoclinic space group) determined by single crystal XRD is shown in Fig. [2](#page-2-0)a. The Fe<sup>II</sup> ions are coordinated sixfold, interconnected via four pseudo square-planar tetracyano-platinate ions, forming 2D metalcyanide-metal sheets. Only two CN- groups of each tetracyanoplatinate are coordinated to the  $Fe<sup>II</sup>$  ions while the two others remain uncoordinated. The coordination sphere of  $Fe<sup>II</sup>$  is completed by the N atoms from the two monodentate, redox-active ligands in an axial configuration. The structure shows two molecules of crystal water forming hydrogen bonds with the uncoordinated CN- moieties of two stacked 2D planes. While this can be seen for the entire family of compounds, compound 3b shows a much higher disorder of the water molecules, enabling us to place them only in partial occupation.

The structure of this family of compounds is the same as the one recently reported by the Real group<sup> $43$ </sup>. The void space in each metal-



Fig. 1 | Redox properties of ligands. a One-electron redox process on the studied R-pbpyCl ligands (R-pbpy<sup>+</sup>= 1-(4-R-phenyl)−4,4'-bipyridinium); b Cyclic voltammetry of  $CH_3$ -pbpy<sup>+</sup> (1) showing the reversible reduction of the bipyridinium unit



(see Supplementary Fig. 6 for R = Br (2), COOCH<sub>3</sub> (3) and NO<sub>2</sub> (4)). [C] =  $10^{-4}$  M in 0.1 M TBAP/CH<sub>3</sub>CN; reference: Ag+|Ag electrode ([Ag<sup>+</sup>]=10<sup>-2</sup> M in CH<sub>3</sub>CN); scan rate: 100 mV⋅s<sup>-1</sup>.

<span id="page-2-0"></span>

Fig. 2 | Crystallographic structure of compound 1b. a unit cell of 1b at 260 K (full crystallographic data for all measured samples are provided in Supplementary Tables); b the structure of 1b is made up of 2D metal-cyanide-metal sheets (here in blue), intercalated by the π−π stacked axial ligands of the sheets above and below. Hydrogen atoms and water molecules were omitted for clarity. All distances are shown in Ångström (Å).

cyanide-metal sheet (see Fig. 2b, in blue) is filled by the π−π stacked axial ligands of the next sheets to either side (see Fig. 2b, in yellow). Taking into account that the entire series presented in this work is isostructural, we provide here experimental proof for the hypothesis proposed in their work $43$ : the presence of a cationic species is responsible for this Hofmann-type structure, with the two anionic non-coordinated CN- units ensuring the overall neutrality of the compound.

Magnetic susceptibility and optical reflectivity measurements for all compounds were performed over 80–325 K (see Fig. 3 for 1-4a, as well as cross for **1-4b**). At room temperature, the compounds are different shades of pink/purple. Upon cooling, a clear and gradual color change to a dark violet/blue is observed in optical reflectivity measurements for all compounds, whereas in magnetic measurements only  $R = CH_3$  and Br show a complete transition from the high spin (HS) to the low spin (LS) state. This difference in the magnetic properties may have its origin in the different electron densities on the ligands. The pyridinium moiety is an electron-deficient group, where the positive charge is delocalized over the aromatic rings. The introduction of an electron-withdrawing group, like  $NO<sub>2</sub>$  or COOCH<sub>3</sub>, leads to the destabilization of the pyridinium unit, whereas the contrary is true for electron-donating groups. In order to prove this hypothesis, a systematic study of the electron donor/acceptor nature of the substituents *vs* SCO should be undertaken, this is, however, out of the scope of the present article.

215K



 $30$  $\chi_{\mathbf{a}^*}$  T (cm $^3$  mol<sup>-1</sup> K) 2.5  $2.0$  $1<sub>5</sub>$  $1.0$  $R = Br$  $0.5$  $0.0$ 50 100 150 200 250 300 350 Temperature (K) 50 45 4.0  $\chi_{a}$ \* T (cm<sup>3</sup> mol<sup>-1</sup> K) 3.5  $3.0$  $2.5$  $2.0$  $1.5$  $1.0$  $R = NO<sub>2</sub>$  $0.5$  $0<sub>0</sub>$  $\mathbf 0$ 50 100 150 200 250 300 350 400 Temperature (K)

Fig. 3 | Magnetic and optical reflectivity measurements of compounds 1-4a.  $\chi_a T$ vs. T ( $\chi_a$  stands for the molar magnetic susceptibility) and variable temperature optical reflectivity in cooling and heating modes for samples of 1-4a

 ${Fe(R\text{-}pbpy^})_2[Ni(CN)_4]_2.2H_2O$  (R = CH<sub>3</sub>, (curve a; red), Br (curve b; blue), COOCH<sub>3</sub> (curve c; yellow) and  $NO<sub>2</sub>$  (curve d; green). The magnetic measurements are shown in the respective colors, the corresponding optical reflectivity curves in grey dots.



Fig. 4 | UV-vis-NIR absorption spectra at 293 K. Compound 1b (purple), {Zn(CH3 **pbpy**<sup>+</sup>)<sub>2</sub>[Pt(CN)<sub>4</sub>]<sub>2</sub>·2H<sub>2</sub>O} (teal) and {Fe(**py**)<sub>2</sub>[Pt(CN)<sub>4</sub>]·2H<sub>2</sub>O} (red).

Fig. [3](#page-2-0): Magnetic and optical reflectivity measurements of compounds 1-4a:  $\chi_a T \nu s T$  ( $\chi_a$  stands for the molar magnetic susceptibility) and variable temperature optical reflectivity in cooling and heating modes for samples of **1-4a** {Fe(**R-pbpy**<sup>+</sup>)<sub>2</sub>[Ni(CN)<sub>4</sub>]<sub>2</sub>·2H<sub>2</sub>O} (R = CH<sub>3</sub>, (curve a; red), Br (curve b; blue), COOCH<sub>3</sub> (curve c; yellow) and  $NO<sub>2</sub>$ (curve d; green). The magnetic measurements are shown in the respective colors, the corresponding optical reflectivity curves in grey dots.

As a color change is characteristic for a spin transition, it is expected that those two phenomena occur at comparable temperatures. Here, however, the optical and magnetic properties are not always correlated. Compounds 1a & 2a show a reversible, gradual change of  $\chi_a T$  with two steps at 200 K and 145 K for R = CH<sub>3</sub> and at 228 K and 18[3](#page-2-0) K for  $R = Br$  (Fig. 3 a, b). For both compounds, the reversible optical transition seems to start first, and is only roughly correlated with the magnetic changes, with the first step leading to a pronounced color change, while the second seems more subtle (vide infra). For compound  $3a$  (R = COOMe) (Fig. [3c](#page-2-0)), the optical change follows the magnetic properties more closely, with a partial SCO at low temperatures. On the contrary, compound  $4a (R = NO<sub>2</sub>)$  (Fig. [3](#page-2-0)d) does not show a SCO in the measured temperature range, however, a pronounced color change is observed anyways. Particularly interesting, it does not only show a gradual color change until the lowest measured temperatures, but also an abrupt transition with a hysteresis loop of 58 K (between 232 K and 290 K). While the overall magnetic behavior of the respective Pt compounds (1-4b, see Supplementary Fig. 8) mirrors the Ni counterparts, the spin transition temperatures lie approximately 10 K higher.

Taking into account the literature on viologen/bipyridiniumbased compounds and the structure of our materials, one hypothesis for the origin of the thermochromism, other than the SCO, is the radical formation within the MOF structure. Indeed as described in literature, this type of compounds can exhibit photo-, thermo- and/or piezo-chromic phenomena due to an electron transfer process originating from donor groups, such as the iron or the non-coordinated CN<sup>−</sup> groups, to the organic ligands $12$ . Thermochromic behavior is observed for many viologen/bipyridinium-based compounds, usually occurring at or above room temperature<sup>14</sup>. In our case, it is possible that the introduction of the bipyridinium unit into a protective/stabilizing framework allows for this phenomenon to occur at lower temperatures. In order to closely examine this hypothesis, we chose two compounds for more detailed analysis: First, {Fe(CH<sub>3</sub>-pbpy<sup>+</sup>)<sub>2</sub>[µ<sub>2</sub>- $M(CN)<sub>4</sub>$ <sub>2</sub>·2H<sub>2</sub>O<sub>3</sub> (1a, b), which allowed us to compare optical and magnetic data with a detailed crystallographic study, and second,

 ${Fe(NO_2\text{-}pbpy^{\dagger})_2}$ [µ<sub>2</sub>-Ni(CN)<sub>4</sub>]<sub>2</sub>·xH<sub>2</sub>O} (4a), in order to gain insight into the non-magnetic transition via an in-depth spectroscopic analysis.

The optical properties of 1a and 1b have been carefully checked by UV-vis-NIR spectroscopy and compared to the parent clathrate compound  ${Fe(pv)}_2[u_4-Pt(CN)_4]$ <sup>2</sup>H<sub>2</sub>O}. At 293 K (HS state), the parent compound presents two main absorption bands centered at 355 and 900 nm, where the near-infrared band is attributed to an MLCT tran-sition (Fig. 4, red curve)<sup>[48](#page-11-0)</sup>. The compound (1b), however, presents an additional absorption band at 535 nm which is rather unusual for HS SCO compounds - it appears when exchanging the innocent pyridine ligand for the redox active  $\mathbf{R}\text{-}\mathbf{p}\mathbf{b}\mathbf{p}\mathbf{v}^+$  unit (Fig. 4, purple curve). In parallel, spectroelectrochemical analysis of the methyl ligand shows that upon reduction at −1.2 V, two new absorption bands emerge at 384 and 545 nm, which are attributed to the reduced, radical form of the ligand (see Supplementary Fig. 7). This indicates, that upon incorporation into the MOF, the ligand might have received an electron leading to the single  $CH_3$ -pbpy' radical, which could explain the strong absorption at 535 nm. It should be noted that this absorption band has been seen for all  ${Fe(\mathbf{R}\text{-}\mathbf{p}\mathbf{b}\mathbf{p}\mathbf{y}^*)}_2[\mu_2\text{-}M(CN)_4]_2\text{-}2H_2O}$  compounds. In addition, isostructural Zn derivatives  $\{Zn(\mathbf{R}\text{-}pbpy^*)_2[\mu_2\text{-}Ni(CN)_4]_2\text{-}2H_2O\}$  were synthesized (see Supplementary Fig. 17 for powder XRD of  $R = CH_3 (1b-Zn)$ and Supplementary Fig. 18 for crystal structure of  $R = Br(2b-Zn)$ ), which, even though they contain the respective redox-active ligand, only show a small shoulder around 500 nm (Fig. 4, teal curve), indicating that the  $Fe<sup>II</sup>$  ion also plays an important role at the origin of this absorption band.

The evolution of the absorbance spectra during the cooling process has been monitored for the parent compound  ${Fe(py)}$  $Ni(CN)<sub>4</sub>$ ]·2H<sub>2</sub>O} (Supplementary Fig. 19), as well as the compounds 1a (see Fig. [5a](#page-4-0), b) and 4a (see Fig. [5c](#page-4-0), d) (check Supplementary Information for compounds 1b (Supplementary Fig. 20), 2a (Supplementary Fig. 21) and 3a (Supplementary Fig. 22)). For 1a, we observe the disappearance of the absorption bands around 1085 nm and 820 nm, as well as the emergence of a shoulder at 575 nm, which is supported by the well-defined isosbestic point until 200 K at 690 nm (see Fig. [5](#page-4-0)a). Typically, if only a SCO is observed in the material, as can be seen for the parent compound  ${Fe(py)_2[\mu_4-Ni(CN)_4] \cdot 2H_2O}$ , the evolution of absorption band intensities (535, 815 and 1000 nm) follows the SCO phenomenon (see Supplementary Fig. 19b). Fig. [5](#page-4-0)b shows the variation of the absorption band intensities with respect to the magnetic and optical properties of 1a, explaining the discrepancy between the two measurements. In contrast to the parent compound, only the evolution of the absorption intensity at 820 nm seems to follow the SCO (see also Supplementary Fig. 20 (1b) and Supplementary Fig. 21 (2a)), the absorption band at 575 nm, however, follows the optical transition, while a contribution at 1085 nm decreases right from the start. This indicates that in addition to the SCO, at least one more thermochromic phenomenon is occurring upon changing the temperature.

Even more intriguing is the behavior of compound 4a, where the observed thermochromism is not related to a spin transition (see Fig. [5](#page-4-0)c, d). Upon cooling, the intensity of the absorption band at 600 nm strongly increases (Fig. [5](#page-4-0)c), which can be superposed with the optical reflectivity measurements. As there is no magnetic transition, no absorption changes were observed around 700–800 nm. For a sideto-side comparison, showing that the color change is fully reversible for the discussed compounds please see Supplementary Fig. 23.

Based on these findings we conclude that the absorption band observed at 580–605 nm corresponds to the optical transition, while the changes around 700 to 800 nm describe the magnetic behavior, thus indicating that besides the SCO at least one more thermochromic phenomenon occurs in each sample. Since the formation of **R-pbpy** radicals gives rise to spectral changes around 500–600 nm as well as around 800–1000 nm (in case of dimer formation)<sup>7[,49](#page-11-0)–52</sup>, it is tempting to associate these events with an ET between the ligand and some donor species (e.g. the iron center or the non-coordinated

<span id="page-4-0"></span>



 $\mathbf{b}$ 

 $*$  T (cm<sup>3</sup> mol<sup>-1</sup> K)

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 $\mathbf{d}$ 

 $\mathbf 0$ 

5

 $\overline{4}$ 

3

 $\overline{2}$ 

3

 $\overline{\mathbf{c}}$ 

**a** compound 1a upon cooling and **b** comparison of the evolution of  $\chi_a$ T and optical reflectivity of 1a and the absorption intensities at selected wavelengths as a function of the temperature; c compound 4a upon cooling and d comparison of the evolution of  $\chi$ <sub>a</sub>T and optical reflectivity of **4a** with the absorption intensity at 600 nm as a function of the temperature upon cooling and heating. The absorption



200



Fig. 6 | Calculated absorption spectra. LS (blue) and HS (red) of {Fe(CH<sub>3</sub>**pbpy**<sup>+</sup>)<sub>2</sub>[ $\mu_2$ -Pt(CN)<sub>4</sub>]<sub>4</sub>}<sup>4</sup> as well as the spectrum of HS {Fe(**CH<sub>3</sub>-pbpy**)<sub>2</sub>[ $\mu_2$ -Pt(CN)<sub>4</sub>]<sub>4</sub>}<sup>6</sup> (yellow).

300

CN<sup>−</sup> groups). To better understand the origin of this SCO-independent thermochromism, TD-DFT calculations and detailed structural analysis (for compound 1b) as well as EPR, Mössbauer, Raman, and Infrared absorption (IR) spectroscopy studies (1a and 4a) have been carried out in the range of 300-80 K.

The electronic excitation calculations were performed within linear-response in time-dependent density-functional theory (LR-TD-DFT) using a simplified model fragment of compound **1b**, based on the crystal structure in the LS state (total electronic spin S=0) and in the HS state  $(S = 2)$ . The low-energy parts of the calculated spectra of LS {Fe(CH<sub>3</sub>-pbpy<sup>+</sup>)<sub>2</sub>[ $\mu_2$ -Pt(CN)<sub>4</sub>]<sub>4</sub>}<sup>4</sup> and HS {Fe(CH<sub>3</sub>-pbpy<sup>+</sup>)<sub>2</sub>[ $\mu_2$ -Pt(CN)<sub>4</sub>]<sub>4</sub>}<sup>4</sup> spectra are dominated by an intense Fe  $\rightarrow$  **CH<sub>3</sub>-pbpy**<sup>+</sup> MLCT transition located at 763 nm and 1185 nm (Fig. 6, blue and red), which correspond to the experimental bands observed at 580 nm and 1065 nm, respectively. The red-shift of the calculated spectra can be ascribed to the approximations underlying the LR-TD-DFT calculations and possibly to the use of a molecular fragment and hence to the neglect of the crystalline environment. The influence of an electron transfer on the optical properties was investigated by performing LR-TD-DFT calculations on the doubly reduced HS fragment (S=3)  ${Fe}$ (CH<sub>3</sub>-pbpy')<sub>2</sub>[ $\mu_2$ -Pt(CN)<sub>4</sub>]<sub>4</sub>]<sup>6</sup> only since the self-consistent-field

303 K

 $303H$ 

 $\chi_a$ \*T

575 K nm

820 K nm 1085 K nm ....optical reflectivity

300

 $\chi$ <sup>\*</sup> T  $600 K nm$  $\cdots$  optical reflectivity

400

ntensity (arb. units)

ntensity (arb. units)

 $333K$ 

350

243 K

 $2431$ 

250

200

Temperature (K)

183 K

150

100

<span id="page-5-0"></span>

**Fig. 7 | Mössbauer spectra of 1a and 4a. 1a** {Fe(**CH<sub>3</sub>-pbpy**\*)<sub>2</sub>[Ni(CN)<sub>4</sub>]<sub>2</sub>·2H<sub>2</sub>O} **a** at 80 K and **b** at 323 K, in comparison to spectra of **4a** {Fe(**NO<sub>2</sub>-pbpy**\*)<sub>2</sub>[Ni(CN)<sub>4</sub>]<sub>2</sub>·2H<sub>2</sub>O} **c** at 80 K and d at 295 K.

calculations on the mono-reduced HS fragment lead to a degenerate electronic state (see Supplementary Discussion). The calculated spectrum of the doubly reduced HS fragment presents two intense bands (Fig. [6](#page-4-0), yellow) that are independent π−π transitions occurring at 796 nm and 663 nm and are centred each on one or the other reduced ligand. The overlap of these intra-ligand absorption bands with the MLCT bands found in the calculated spectra of LS and HS  ${Fe}$ (CH<sub>3</sub>-pbpy<sup>+</sup>)<sub>2</sub>[ $\mu$ <sub>2</sub>-Pt(CN)<sub>4</sub>]<sub>4</sub>}<sup>4</sup> supports the conclusion drawn from the experiments that the temperature dependence of the absorption intensities is likely also due to the presence of reduced  $CH<sub>3</sub>$ -pbpy ligands at elevated temperatures.

EPR does not show a signal characteristic for a radical, which is, however, to be expected in the case of dimer formation<sup>7[,49](#page-11-0)-52</sup>. This hypothesis is also in agreement with the observation of an absorption band near 800–1000 nm, characteristic of bipyridinium dimer species<sup>52</sup>, and with the observation of  $π$ -π interactions between the ligands (vide infra).

 $57$ Fe Mössbauer spectroscopy is a useful technique to determine the spin and oxidation states of iron. Indeed, in case of an ET event from the Fe<sup>II</sup> central ion to the ligand, a characteristic signature of Fe<sup>III</sup> might be visible via Mössbauer studies. Thus, Mössbauer spectra have been recorded for compounds 1-4a in the range of 300–80 K (see Supplementary Fig. 24–27). In Fig. 7a, b the spectra at 80 K and 323 K are shown respectively for 1a (see Supplementary Fig. 24 for additional temperatures). At 323 K an asymmetric doublet is observed, which can

be clearly assigned to a  $Fe<sup>II</sup>_{HS}$  species based on the values of isomer shift ( $\delta^{CS}$  = 1.005(17) mm s<sup>-1</sup>) and quadrupole splitting ( $\Delta E_{Q}$  = 0.987(35) mm s<sup>-1</sup>). Upon cooling, the intensity of this doublet gradually decreases whereas a new doublet emerges, and finally, at 80 K, the sample shows a fully symmetric  $Fe^{II}$ <sub>LS</sub> doublet ( $\delta^{CS}$  = 0.4674(40) mm s<sup>-1</sup> and  $\Delta E_{\rm Q}$  = 0.2353(64) mm s<sup>-1</sup>). These observations are characteristic of a spin state switching between  $Fe<sup>H</sup>_{HS}$  (S = 2) and  $Fe<sup>H</sup>_{LS}$  (S = 0), and can be correlated to the magnetic data. The asymmetry of a quadrupole-split Mössbauer line at high temperatures remains, however, intriguing. Such line asymmetry may have different origins, including preferential crystalline orientation (texture), lattice vibrational anisotropy (Goldanskii-Karyagin effect), a residual  $Fe^{II}_{LS}$  species, or a relaxational line broadening<sup>53</sup>. Nevertheless, based on the temperature dependence of the spectrum, characterized by a well-defined shoulder at higher temperatures and a symmetrical doublet at the lowest temperatures, the best fit of the spectra is obtained by assuming the existence of a second iron species ( $\delta^{CS}$  = 0.24(10) mm s<sup>-1</sup> and  $\Delta E_{\rm Q}$  = 0.55(17) mm s<sup>-1</sup>) with a population of ca.  $29 \pm 8$  % at 323 K. This species might be ascribed to a ferric ion, possibly  $Fe^{III}$ <sub>HS</sub>, and shows a notable temperature dependence, indicating that it does not stem from the accidental oxidation of the samples. The Mössbauer results thus support the UV-visible data, showing that a part of the iron ions participates in an ET towards the ligands when increasing the temperature – in partial overlap with the SCO phenomenon. One shall also note that there is only one SCO active iron site (confirmed both by Mössbauer and XRD

<span id="page-6-0"></span>



Fig. 8 | Vibrational spectroscopy analysis of compounds 1a and 4a. a Raman spectra (acquired with a 785 nm laser) of 1a recorded at 323 K (red), 193 K (blue), and 93 K (yellow); b Raman spectra (acquired with a 785 nm laser) of 4a recorded at

283 K (red), 243 K (blue) and 133 K (yellow); c temperature dependent Infrared absorption spectrum of compound 1a and d temperature dependent Infrared absorption spectrum of compound 4a.

measurements). This means that the two-step SCO observed in mag-netism (see Fig. [3](#page-2-0)a, b) is not linked to the presence of two distinct  $Fe<sup>II</sup>$ sites, but could arise due to a coupling of different physical phenomena. A plausible hypothesis is a coupling between the SCO and ET events.

Figs [7c](#page-5-0), d show the Mössbauer absorption spectra for compound 4a at 80 K and 295 K respectively (see Supplementary Fig. 26 for additional temperatures). Even though we were able to improve spectral quality by enriching sample  $4a$  with  $57Fe$ , it is difficult to determine a unique fit due to the rather large doublet present in the spectra. At ambient temperature, a symmetric doublet is observed, consisting of at least two Fe<sup>II</sup> HS species (Fe<sup>II</sup> HS (1)  $\delta$ <sup>CS</sup> = 1.114(20) mm s<sup>-1</sup> and  $\Delta E_O = 1.493(85)$  mm s<sup>-1</sup> and (2)  $\delta$ <sup>CS</sup> = 1.1330(55) mm s<sup>-1</sup> and  $\Delta E_{\rm Q}$  = 1.016(13) mm s<sup>-1</sup>). Upon cooling we see the emergence of a new  $Fe<sup>II</sup>$  HS species with a large quadrupole splitting, reaching up to 20 % population at 80 K (Fe<sup>II</sup> HS (3)  $\delta^{CS} = 1.2290(88)$  mm s<sup>-1</sup> and  $\Delta E_{\rm Q}$  = 2.800(37) mm s<sup>-1</sup>). This indicates that the coordination environment of part of the iron centers presents changes with temperature, which might stem from an electron transfer to the ligand and explain the color change observed for this sample. The asymmetric nature of the spectra at low temperatures is partially attributed to the increase of a fourth Fe species ( $\delta^{\text{CS}}$  = 0.5326(76) mm s<sup>−1</sup> and  $\Delta E_{\text{Q}}$  = 0.302(11) mm s<sup>−1</sup>), that might correspond to a small LS fraction or a Fe<sup>III</sup> species. Since we do not observe a SCO in compound 4a, we believe that the latter is more likely.

As both the SCO and ET phenomena should be reflected by changes in vibrational spectroscopy, variable temperature Raman and Infrared (IR) spectra of the samples were acquired (see Fig. 8). Raman microscopy measurements of 1a (see Fig. 8a) reveal new peaks during the cooling process (e.g. at  $685 \text{ cm}^{-1}$ ) and, in addition, the intensity ratio between peaks located around 1612 cm<sup>-1</sup> changes. These peaks are attributed to the stretching and deformation vibrations of the C-H and C-N bonds of the ligand. Moreover, a shift of the CN modes to higher wavenumbers, caused by the SCO, can be observed (see Supplementary Fig. 28). Compound 4a (see Fig. 8b) shows similar behavior with new peaks around 659 and 1275 cm<sup>-1</sup> and an intensity ratio change between peaks located around 1620 cm<sup>-1</sup>. As for compound 1a, it shows two distinct C-N stretching modes, since only two of the four CN<sup>−</sup> moieties are coordinated to the Fe<sup>II</sup>, however, as expected in this case, no CN mode shift can be detected.

The analysis of the Raman spectral changes is complicated by the strong thermochromism of the samples, potentially leading to the resonance enhancement of certain Raman modes. IR absorption studies (see Fig. 8c, d), on the other hand, enabled us to follow the temperature-induced changes of vibrational modes with less ambiguity. Importantly, we observed significant changes in the out-of-plane CH bending regions (both fundamental and overtone) of the aromatic ligands both in the SCO active and inactive derivatives (see Supplementary Figs. 29 and 30, for region of 900 to 1200 cm<sup>-1</sup> of compound 1a and 4a, respectively). This behavior strongly contrasts with that of the parent clathrate compound  ${Fe(py)_2[\mu_4-Ni(CN)_4] \cdot 2H_2O}$ , which displays only slight shifts of vibrational modes associated with the pyridine ligand<sup>54</sup>. Since the compounds are isostructural at each temperature (no symmetry change) we can thus infer a significant variation in the structure of the  $\mathbf{R}\text{-}\mathbf{p}\mathbf{b}\mathbf{p}\mathbf{y}^*$  ligands upon temperature changes. Indeed, the uncoordinated ν(C-N) modes display significant changes as

<span id="page-7-0"></span>

Fig. 9 | In-detail crystallographic analysis of compound 1b. a side view of two πstacked units of 1b at 290 K, indicating where interactions and structural changes are observed in temperature. Fe-N distances are marked in light blue, the Fe-N-C angle in purple, the π−π stacking and bipyridinium-NC distances in red and green, respectively, and torsion angles in orange. Water molecules are omitted for clarity.

b Evolution of the distance Fe1-N1 and the torsion angle of the flat bipyridinium unit (C011-C019-C01B-C00U) upon cooling of 1b compared to its magnetic behavior, error bars designate standard deviation; c selection of metal-to-ligand bond lengths  $(A)$  and torsion angles  $(°)$  for **1b** at 150 K and 290 K.

a function of the temperature as well. Compound 1a (Fig. [8](#page-6-0)c) presents two asymmetric doublets at 2159 and 2135 cm<sup>−</sup><sup>1</sup> attributed to the two different C-N modes (coordinated and uncoordinated, respectively). When increasing the temperature, a new peak emerges at  $2120 \text{ cm}^{-1}$  in the spectral region attributed to the uncoordinated CN<sup>-</sup> units. For compound 4a a similar behavior is observed with a shoulder emerging in the same position (see Fig. [8](#page-6-0)d). These observations can be related to an ET phenomenon between the CN<sup>-</sup> group, which could act as an electron donor, and the bipyridinium ligands. Similar arguments have been proposed in the paper of Guo et al. for the origin of the ET observed in their compound<sup>12</sup>.

As mentioned in the introduction, the occurrence of ET phenomena in compounds containing bipyridinium derivatives is associated with several subtle structural features. According to the Marcus ET theory, the shorter the distance between an electron donor and acceptor the faster the ET will occur, the maximum distance possible being around 4 Å. The ET is often related to the presence of hydrogen bonds and favored by π−π interactions in the crystal packing. In addition, the transfer is more likely to occur if the interannular torsion angle of the pyridinium units is negligible $14$ . To explore the structural details, a thorough crystallographic study of the single crystal 1b  ${Fe}$ (CH<sub>3</sub>-pbpy<sup>+</sup>)<sub>2</sub>[µ<sub>2</sub>-Pt(CN)<sub>4</sub>]<sub>2</sub>·2H<sub>2</sub>O} was undertaken at various temperatures. At 300 K, the average Fe–N lengths for the cyano and viologen units vary between 2.15 and 2.22 Å (see Fig. 9c), which is in close agreement with a Fe<sup>II</sup> ion in the HS state. Interestingly, the axial  $CH_3$ pbpy<sup>+</sup> ligands are non-equivalent in regards to the interannular angle of the bipyridinium unit: while one shows a torsion angle of around 44°, the other unit is almost fully planar (see Fig. 9a, c). Moreover, very short distances of less than 3 Å can be observed between the uncoordinated CN- units and the bipyridinium units of the axial ligands. In addition, the uncoordinated CN<sup>−</sup> groups show a clear structural disorder at higher temperature (280 K and 290 K), confirming the observations made in IR spectroscopy. Furthermore, π−π interactions between two  $CH_3$ -pbpy<sup>+</sup> ligands are present, possibly explaining the EPR silent behavior of the compounds. All four phenomena are in line with the aforementioned requirements for an ET at ambient



Fig. 10 | Cyclic voltammetry and Raman spectroscopy of compound 1a. a Cyclic voltammetry of la versus Ag<sup>+</sup>|AgCl in the solid state. 5 mg of the target material were dispersed in 500 µL of a 4% PVA solution in water and vigorously sonicated, then 125 µL of a 4% BORAX solution in water were added, forming a gel that was

then applied onto a glassy carbon electrode (3 mm diameter). The scan rate was 50 mV s<sup>-1</sup> in 1 M KCl; **b** in-situ Raman spectroscopy measurements of the **1a**, showing the pristine clathrate (red) after the reduction (blue) and after re-oxidation (yellow). The dashed lines highlight the new peaks of the reduced compound.

temperature within our structure and would suggest an electron transfer from the uncoordinated CN- groups to the axial ligand, in agreement with the evolution of the IR absorption bands discussed above.

Upon cooling, significant changes can be observed in the crystal structure. At 150 K the spin transition is virtually complete as indicated by the typical shortening of the Fe-N distances for both the cyano and pyridine units to 1.945 Å and 2.013 Å, respectively (see Fig. [9b](#page-7-0), c). It is important to mention that the bipyridinium unit showing a fully planar interannular angle is strongly affected by temperature, reaching a more twisted state with an angle of 12–14° when cooling to 150 K. As a thermo-induced ET is much more likely in flat bipyridinium units compared to twisted ones, this confirms that the process will more likely take place at (or above) ambient temperature. In addition, it is interesting to notice that the distances between the uncoordinated CNand bipyridinium units do not show a large variation in temperature, even though we observe a shortening of the Fe-N bonds occurring upon the spin transition. This is another indication for the coupling between the uncoordinated CN<sup>-</sup> and bipyridinium units. Overall, the planar configuration of the bipyridinium unit at 300 K combined with the short distances to the uncoordinated CN moiety, which likely acts as an electron donor, is perfectly in line with the hypothesis of an intramolecular ET within the structure.

The multifunctionality of the compounds induced by the presence of a redox-active ligand, has been highlighted through cyclic voltammetry and spectro-electrochemical studies (see Fig. 10). There are several examples in the literature concerning the chemical reduction of Hofmann-type clathrates in order to modulate the SCO properties. Ohba et al.<sup>55</sup> conducted a post synthetic oxidative addition of iodine to their compound, which coordinates to the Pt ions, oxidizing them from  $Pt^{\text{II}}$  to  $Pt^{\text{IV}}$ . Depending on the iodine content and due to pore size restrictions, they were able to oxidize up to half of all Pt ions, thereby modulating the transition temperature of their compound in a range of 300 to 400 K, while maintaining the cooperative transition. Similarly, Zuo et al.<sup>56</sup> doped a tetra(4-pyridyl)tetrathiafulvalene based  $Fe<sup>II</sup>$  MOF with  $I_2$ , thereby modifying its magnetic behavior and increasing the MOFs conductivity. Tong et al. $57$  reported a cationic metal organic framework with larger pores, where the chemical oxidation by coordination of iodine or bromine was possible for all  $Au<sup>II</sup>$ sites, thus modifying the SCO behavior from a hysteretic one-step transition to a three- or two-step transition, respectively. They show that after the subsequent reduction of the compound using ascorbic acid, the one-step hysteretic transition can be reinstated, however, with permanent changes in spectroscopic and SCO properties. Coronado et al[.58](#page-11-0) have reported a SCO active polyoxometalate (POM)

compound, that can undergo a chemical or electrochemical reduction due to the redox-active linkers employed. They show this reduction to be reversible, the magnetic properties of the chemically re-oxidized compound are, however, modified. Recently, a review focusing on SCO and valence tautomerism conductors has been published, discussing synthesis strategies, magnetic and conductive properties as well as the influence of the materials inherent bistability on their conductivity<sup>[59](#page-11-0)</sup>. Interesting concepts are the co-crystallization of a bistable matrix with a conductive unit (often a redox-active, planar molecule) – here, SCO materials offer interesting prospects due to the volume change seen upon the spin transition.

Since the Hofmann clathrate derivatives presented in this work are not soluble, their electrochemical redox properties were studied in the solid state, discussed here for the  $R=CH_3$  derivative. **1a** shows a reduction potential of  $E_{1/2} = -0.765$  V vs. Ag<sup>+</sup>|AgCl (see Fig. 10a), which is ascribed to the one-electron reduction of the pyridinium unit, in line with previous reports in the literature on the electrochemical behavior of bipyridinium derivatives $46,47$ . The quasi-reversible cyclic voltammogram shows a higher current intensity for the anodic wave, which might be related to the dissolution of the compound. In addition, compound 1a was investigated spectro-electrochemically at a negative potential. Fig. 10b shows the Raman spectra of 1a before and after reduction, followed by a re-oxidation. After the reduction at −1.0 V vs. Ag<sup>+</sup>|AgCl, we remark a complete modification of the Raman spectra with the emergence of several new peaks (e.g. at 995, 1190, and 1537 cm<sup>−</sup><sup>1</sup> ) in the region attributed to the stretching vibrations of the ligand. These modifications are ascribed to the one-electron reduction of the bipyridinium ligand within the MOF structure. The re-oxidation shows the reversibility of the system as the spectrum of the pristine compound is fully recovered. This result contrasts with the abovementioned studies carried out on the chemical reduction or oxidation of similar compounds and proves electrochemistry to be a useful method to obtain multifunctional materials. It shows that this family of compounds is able to respond to multiple external stimuli such as temperature or an applied potential in order to attain several stable and reversible states, allowing us to classify these materials as multifunctional.

In conclusion, we have shown that the rational design of a series of non-innocent ligands for Hofmann-type clathrates affords a family of redox-active SCO compounds,  ${Fe(\mathbf{R}\text{-} \mathbf{pbpy}^*)_2[\mu_2 \text{-}M(CN)_4]_2 \cdot 2H_2O}$  (R =  $CH_3$ , Br, COOCH<sub>3</sub> or NO<sub>2</sub>; M = Ni or Pt), displaying an unusual structural arrangement due to the cationic nature of the ligand. The investigation of the optical properties of the compounds over a temperature range of 100–300 K reveals their thermo-chromic character which cannot be solely ascribed to the SCO phenomenon. Indeed, the combination of

spectroscopic, crystallographic, and theoretical findings points to the occurrence of at least one additional electron transfer process and the associated formation of radical dimers within the structure. From Mössbauer and UV visible data, it is clear that the iron ion plays an important role in the ET properties. In addition, vibrational spectroscopies show strong changes in the stretching modes of the uncoordinated CN<sup>−</sup> group upon cooling. This result, and the observed short distances of 3 Å between the bipyridinium unit and the uncoordinated CN<sup>−</sup> moieties indicate the latter as the most likely origin of the ET. In addition, the presence of  $π$ -π interactions between the bipyridinium ligands, which stabilize the formed dimers, as well as the small interannular torsion angle observed for the latter, satisfy the structural requirements for an ET to occur. Importantly, we have shown that this family of compounds is able to undergo a quasi-reversible electrochemical redox process that can be followed via in situ Raman spectroscopy. This last result confirms the multi-stability and multiaddressability of our compounds, thus opening the door to a new class of switchable materials.

# Methods

### Materials

All purchased chemicals were used as received:  $Fe(BF<sub>4</sub>)<sub>2</sub>·6H<sub>2</sub>O$ : Iron(II) tetrafluoroborate,  $K_2[M(CN)_4]$ : Potassium tetracyanometallate (M = Ni or Pt), 4,4'-bipyridine, 2,4-dinitrochlorobenzene, trimethylamine, methyl-4 aminobenzoate, 4-methylaniline, 4-bromoaniline, 4-nitroaniline.

### Synthesis

Ligands were synthesized in two steps: First, 20 mmol (3.12 g) 4,4' bipyridine and 20 mmol (4.05 g) 2,4-dinitrochlorobenzene are dissolved in 50 ml acetone and stirred at reflux for 12 h, cooled, then the precipitate that formed during reaction (di-NO<sub>2</sub>-pbpyCl) was filtered and washed multiple times with acetone (Yield: 77%). Second, 3 equivalents of a 4-R-aniline and 1 equivalent of  $di$ -NO<sub>2</sub>-pbpyCl were dissolved in 15 ml of ethanol abs. stirred at reflux for 4 h, cooled, and then, 20 ml dist.  $H_2O$  was added to remove the byproduct as precipitate. All solvent was evaporated and the crude product dissolved in as little methanol as possible, then, the ligand R-pbpyCl was precipitated in diethyl ether and washed multiple times with diethylether to remove impurities. Please check the Supplementary Methods section for a more detailed description and the characterisation (<sup>1</sup>H-NMR and mass spectrometry results) of all synthesized ligands. Clathrates were synthesized with 0.5 mmol of the respective ligand  $(L = CH<sub>3</sub>$ pbpyCl, COOCH<sub>3</sub>-pbpyCl, NO<sub>2</sub>-pbpyCl or Br-pbpyCl), that are dissolved in 10 mL 50/50 v% EtOH/H<sub>2</sub>O dist., then 0.25 mmol of Fe(BF<sub>4</sub>)<sub>2</sub> · 6 H2O are added with a small amount of ascorbic acid. Once dissolved, 0.5 mmol of  $K_2[Ni(CN)_4]$  x H<sub>2</sub>O or  $K_2[Pt(CN)_4]$  are added, and purple precipitates form immediately. The mixture is stirred for 1 h, filtered and the precipitate is washed several times with EtOH/H<sub>2</sub>O dist., then dried in the oven. The purity of the microcrystalline powders was investigated via CHN and Inductively Coupled Plasma (ICP) analysis as well as powder X-ray diffraction (please see Supplementary Methods and Supplementary Figs.). Crystals were grown with 0.35 mmol of the respective ligand  $(L = CH_3$ -pbpyCl, COOCH<sub>3</sub>-pbpyCl or BrpbpyCl), that are dissolved in 8 mL 50/50 v% EtOH/H2O dist. and filled into an H-tube, then 0.1 mmol of  $Fe(BF_4)_2 \cdot 6 H_2O$  in 0.5 mL 50/50 v % EtOH/H<sub>2</sub>O dist. are added on one side and 0.2 mmol of  $K_2[Pt(CN)_4]$  in 0.5 mL 50/50 v% EtOH/H<sub>2</sub>O dist. are added on the other side. After several days/weeks, red crystals are formed and characterized via single-crystal X-ray diffraction.

### **Measurements**

Elemental analyses were performed using a Perkin Elmer 2400 series II Instrument for CHN analysis and ICAP 6300 Thermo RF Generator with CID detector for Inductively Coupled Plasma analysis. Mass Spectrometry was performed on a ThermoScientific LCQ Fleet mass spectrometer with ion trap in electrospray ionization mode. Nuclear Magnetic Resonance (NMR), <sup>1</sup>H-NMR spectra were obtained at 298 K in DMSO-d6 as an internal reference and were recorded on a Bruker Avance 300 or Bruker Avance 400. Chemical shifts are reported in parts per million (ppm). Optical reflectivity measurements were carried out on a Linkam THMS 600 heating-cooling stage for microscopy, where a few mg of sample were placed and flattened. The samples were heated to 323 K and the stage was purged under dry N2 flow for 30 min. Then the samples were cooled to 93 K and heated back up to 323 K at a speed of 5 °C/min for two cycles. Pictures were taken every 12 seconds using a Motic stereomicroscope equipped with a color camera and graphically analyzed using the ImageG software. UV-vis-NIR spectra of powders were obtained on a Perkin Elmer LAMBDA 950 Spectrophotometer with an Integrating Sphere and a spectral range of 200 to 2500 nm in diffuse scattering mode. For temperature-dependent measurements, the samples were placed inside a Linkam stage (see above) and thermalized for 2 min at the respective temperature before taking a UV-vis-NIR spectra. Raman spectra were measured on an Xplora (Horiba) Raman micro-spectrometer equipped with an edge filter and a diode laser (785 nm, 0.1 mW). The laser beam was focused on the sample by an  $\times$ 50 objective (numerical aperture = 0.5), which was also used to collect the scattered photons. For spectral calibration, a pure Si wafer was used. Spectra were recorded with an acquisition time of  $6 \times 10$  s from 50 to 2300 cm<sup>-1</sup>. The measurements were carried out using a Linkam heating stage for microscopy (see above). Spectra were recorded each 10 °C after the temperature had stabilized. FTIR spectra were recorded on a PerkinElmer Frontier FTIR/FIR instrument equipped with a Spotlight 150 IR microscope in a range of 4000 to 600 cm<sup>-1</sup> at a resolution of 0.125 cm<sup>-1</sup> and 16 scans. The measurements were carried out using a Linkam heating stage for microscopy (see above) using ZnSe windows. Spectra were recorded each 10 °C after the temperature had stabilized. 57Fe Mössbauer spectra were recorded using a conventional constant-acceleration-type spectrometer equipped with a 57Co source and a liquid nitrogen cryostat. Least-squares fittings of the Mössbauer spectra were carried out with the assumption of Lorentzian line shapes using the Recoil software. The isomer shift values are given with respect to metallic iron at room temperature. Variable-temperature magnetic susceptibility data were obtained at cooling and heating rates of 2 or 4 K min<sup>−</sup><sup>1</sup> under a field of 1 kOe using a Quantum Design MPMS magnetometer. Electrochemical measurements were conducted in a glovebox with a standard one-compartment, three-electrode electrochemical cell using a BioLogic SP300 potentiostat. Experiments in solution were carried out in anhydrous  $CH<sub>3</sub>CN$  containing tetra-n-butylammonium perchlorate (0.1 M) as the supporting electrolyte. The solution concentrations used for the electrochemical studies were typically 10<sup>−</sup><sup>4</sup> M. The counter electrode was a platinum wire, and an ALS Co. Ltd. Ag<sup>+</sup>|Ag electrode ([Ag+] =  $10^{-2}$ M in CH3CN) served as the reference. This electrode was separated from the solution by a bridging compartment. For CV measurements, the working electrode was a 3 mm diameter vitreous carbon disk, which was polished with 1 μm diamond paste before each recording. UV-visible spectroelectrochemical analyses were carried out using a diode array spectrophotometer equipped with a dip probe (10 mm) connected to the spectrophotometer through an optical fiber. Measurements on solid-state materials were carried out in 1 M KCl (aq), using an aqueous reference electrode ALS Co. Ltd. Ag<sup>+</sup>|AgCl (3 M NaCl) and a Pt wire as counter electrode. 5 mg of the target material were dispersed in 500  $\mu$ L of a 4% PVA solution in water and vigorously sonicated, then 125 µL of a 4 % BORAX solution in water were added, forming a gel that was then applied onto a glassy carbon electrode (3 mm diameter). The scan rate was  $100 \text{ mV s}^{-1}$ .

#### Powder and Single-crystal X-ray crystallographic studies

The powder X-ray diffraction (XRD) analyzes were carried out at room temperature on a Rigaku Miniflex600 benchtop powder

<span id="page-10-0"></span>diffractometer in a theta/2 theta geometry using a Cu anode (K-Alpha1 [Å] 1.54060), a fast 1D detector with energy selection, and a 6-position sample changer. Samples were measured in a range of 2theta from 5 to  $60^{\circ}$ , every 0.02°, with a speed of  $5^{\circ}$  per minute. For single crystal XRD, the crystals were kept in their mother liquor until they were dipped into perfluoropolyether oil and their structure was determined. The chosen crystals were mounted on a Mitegen micromount. Data were collected at different temperatures (from 100 K to 300 K) on a XtaLAB Synergy (Rigaku) diffractometer using either a Cu-Kα radiation  $(\lambda = 1.54184 \text{ Å})$  or a Mo-Kα radiation  $(\lambda = 0.71073 \text{ Å})$  micro-source, equipped with an Oxford Cryosystems Cooler Device. The structures have been solved using the new dualspace algorithm program  $SHELXT<sup>60</sup>$  and refined by means of least-squares procedures using SHELXL-2018<sup>[60](#page-11-0)</sup> program included either in the software package WinGX $61$  version 1.639 or the package OLEX2 $62$ . The Atomic Scattering Factors were taken from International Tables for X-Ray Crystallography<sup>[63](#page-11-0)</sup>. Hydrogen atoms were placed geometrically and refined using a riding model, except for the hydrogens of water molecules which were located by Fourier Differences. All nonhydrogen atoms were anisotropically refined. Ellipsoid plots in the figures of the crystallography section were generated using the software ORTEP-3<sup>64</sup>. The crystal structures have been deposited at the Cambridge Crystallographic Data Centre and allocated the deposition numbers 2321396-2321412, 2327738, 2327739, and 2355273.

#### Calculations of electron absorption spectra

All calculations have been performed with the ADF program $65,66$  $65,66$  using the statistical average of orbital potentials (SAOP) exchangecorrelation potential and including scalar-relativistic effects within the zero-order regular approximation  $(ZORA)^{67,68}$ . The atoms were described by relativistic all-electron Slater-type basis sets from the ADF program[69](#page-12-0): the TZP basis set of triple-ζ polarized quality was employed for the Fe and Pt atoms, and the DZP basis set of double-ζ polarized quality for the N, C, and H atoms. The fragments were solvated in water using the conductor-like screening model of solvation  $(COSMO)^{70}$ . The electronic absorption properties have been studied by performing electronic excitation calculations within linear response in timedependent density functional theory  $(LR\text{-}TDDFT)^{71,72}$  and the Tamm-Dancoff (TDA) approximation<sup>73</sup>. The analysis of the predicted transitions was done in terms of transition between occupied and virtual natural transition orbitals (NTO) $74$ . The calculations were run restricted for the LS state and unrestricted for the considered HS state with the projection of the total spin moment S along the z-axis constrained to  $MS = +S$ .

### Data availability

The X-ray crystallographic data for samples 1b (from 150 K to 290 K, one structure every 10 K ((CCDC-2321398 to (CCDC-2321412)), 2b (CCDC-2327738 (100 K), CCDC-2327739 (300 K)) and 3b (CCDC-2321396 (100 K), CCDC-2321397 (300 K)), as well as 2b-Zn (CCDC-2355273 (300 K)) generated in this study have been deposited at the Cambridge Crystallographic Data Center (CCDC). These data can be obtained free of charge from The Cambridge Crystallographic Data Center via [https://www.ccdc.cam.ac.uk/structures/?.](https://www.ccdc.cam.ac.uk/structures/) The corresponding tables are provided in the Supplementary Information file. We declare that the data supporting the findings of this study are available within the article and its Supplementary Information files. All relevant source data are available from the corresponding author upon request.

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# Author contributions

LG&SC (experimental work and manuscript), DP&LV (crystallographic studies), LMLD (theoretical calculations), AR (magnetic measurements), and GM&AB (manuscript).

## Competing interests

The authors declare no competing interests.

# Additional information

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