

### Showcasing research from the group of Prof. Peter Weinberger (Institute of Applied Synthetic Chemistry, TU Wien, Austria).

Bifunctional Fe(11) spin crossover-complexes based on  $\omega$ -(1*H*-tetrazol-1-yl) carboxylic acids

Supramolecular cooperativity resulting in more abrupt spin state transitions is governed by a subtle interplay of ligand functionalization with carboxylic acid moieties, spacer length, weak-coordinating anion and solvate molecules.



See Danny Müller, Peter Weinberger *et al., Dalton Trans.,* 2020, **49**, 17183.



rsc.li/dalton Registered charity number: 207890

# Dalton Transactions

# PAPER

Check for updates

Cite this: Dalton Trans., 2020, 49, 17183

Bifunctional Fe(II) spin crossover-complexes based on  $\omega$ -(1*H*-tetrazol-1-yl) carboxylic acids<sup>†</sup>

Willi Zeni,<sup>a</sup> Marco Seifried,<sup>a</sup> Christian Knoll, <sup>b</sup> <sup>a</sup> Jan M. Welch,<sup>b</sup> Gerald Giester,<sup>c</sup> Berthold Stöger,<sup>d</sup> Werner Artner,<sup>d</sup> Michael Reissner,<sup>e</sup> Danny Müller <sup>b</sup> \*<sup>a</sup> and Peter Weinberger <sup>b</sup> \*<sup>a</sup>

To increase the supramolecular cooperativity in Fe(II) spin crossover materials based on N1-substituted tetrazoles, a series of  $\omega$ -(1*H*-tetrazol-1-yl) carboxylic acids with chain-lengths of C<sub>2</sub>-C<sub>4</sub> were synthesized. Structural characterization confirmed the formation of a strong hydrogen-bond network, responsible for enhanced cooperativity in the materials and thus largely complete spin-state transitions for the ligands with chain lengths of C<sub>2</sub> and C<sub>4</sub>. To complement the structural and magnetic investigation, electronic spectroscopy was used to investigate the spin-state transition. An initial attempt to utilize the bifunctional coordination ability of the  $\omega$ -(1*H*-tetrazol-1-yl) carboxylic acids for preparation of mixed-metallic 3d-4f coordination polymers resulted in a novel one-dimensional gadolinium-oxo chain system with the  $\omega$ -(1*H*-tetrazol-1-yl) carboxylic acids not polymers by the spin-state transition.

Received 23rd September 2020, Accepted 26th October 2020

DOI: 10.1039/d0dt03315d

rsc.li/dalton

## Introduction

First row transition metals with  $3d^4-3d^7$  electron configurations allow for population of their 3d-orbitals with either a maximum, or a minimum of paired electrons. Depending on the coordinative environment around the metal centre, the energetic difference between these electron configurations (high-spin (HS) and low-spin (LS) state) may be small enough to be governed by an external stimulus such as temperature,<sup>1,2</sup> pressure,<sup>1-3</sup> light<sup>4</sup> or external electric field,<sup>5-8</sup> among others.<sup>1</sup> This phenomenon was first observed in the 1930s by Italian researchers, working on iron(III) *N*,*N*-dialkyldithiocarbamates and has since been known as spin crossover (SCO), or spin state transition.<sup>9–11</sup>

The intrinsic correlation between the bistability of SCO materials and several of their physical properties makes them

<sup>a</sup>Institute of Applied Synthetic Chemistry, TU Wien, Getreidemarkt 9/163-AC, 1060 Vienna, Austria. E-mail: danny.mueller@tuwien.ac.at, appealing candidates for novel miniaturized sensors,<sup>12,13</sup> memory/storage devices,<sup>14</sup> and a key technology in spintronics.<sup>15</sup> Today, SCO materials are primarily investigated by an academic bottom-up approach, resulting in highly specialized materials for devices on a nanoscale. Molecular actuators,<sup>16</sup> their utilization in quenching and affecting luminescence and fluorescence,<sup>17,18</sup> their sensing capacity in form of porous switchable materials,<sup>12,19</sup> or their combination with chirality<sup>20–22</sup> have recently been reported.

Since the occurrence of SCO under ambient conditions is very sensitive to the ligand field, it is strongly affected by the nature of the ligand system, the presence or absence of solvates<sup>23</sup> and counter anions applied.<sup>24</sup> (Substituted) azole ligands, including N1-substituted tetrazoles, are a promising platform for systematic development of SCO-materials.<sup>1</sup>

Homoleptic hexa-coordinated Fe(II)-complexes based on N1substituted tetrazoles are often spin switchable, as coordination of Fe(II) by tetrazoles results in an appropriate ligand field strength. In this case, Fe(II) switches between a paramagnetic HS state with S = 2 and a diamagnetic LS state with S = 0.

In previous work, we have emphasized fine-tuning of the ligands' geometry and flexibility, shaping cooperativity and transition temperature on a molecular scale,<sup>25,26</sup> as well as post functionalization of the ligand backbone.<sup>27</sup> Therefore, in this work, a carboxylic acid moiety has been introduced to the ligand system, allowing for the investigation of the SCO in [Fe ( $\omega$ -(1*H*-tetrazol-1-yl) carboxylic acid)<sub>6</sub>]<sup>2+</sup>-cations. Alkylcarboxylic acid substituted tetrazoles provide three useful features: On a molecular level the COOH-group should establish a network of hydrogen bonds, increasing cooperativity and governing the



View Article Online View Journal | View Issue

peter.e163.weinberger@tuwien.ac.at

<sup>&</sup>lt;sup>b</sup>Center for Labelling and Isotope Production, TRIGA Center Atominstitut, TU Wien, Stadionallee 2, 1020 Vienna, Austria

<sup>&</sup>lt;sup>c</sup>Department of Mineralogy and Crystallography, University of Vienna, Althanstraße 14 (UZA 2), 1090 Vienna, Austria

<sup>&</sup>lt;sup>d</sup>X-Ray Center, TU Wien, Getreidemarkt 9, 1060 Vienna, Austria

<sup>&</sup>lt;sup>e</sup>Institute of Solid State Physics, TU Wien, Wiedner Hauptstraße 8-10/138, 1040 Vienna, Austria

<sup>†</sup>Electronic supplementary information (ESI) available. CCDC 2024140 (4, HS), 2024142 (4, LS), 2024146 (6, HS), 2024147 (6, LS), 2024143 (7, HS), 2024141 (7, LS), 2024144 (9, HS), 2024145 (9, LS) and 2024832 (10). For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/d0dt03315d

abruptness of the spin state transition,<sup>28–30</sup> the COOH-group may act as an additional ligand allowing for the formation of multi-metallic networks with a second transition metal, or rare-earth element and, finally, the COOH termination of the ligand may allow for deposition on oxide surfaces.

In this contribution, we report the  $\omega$ -(1*H*-tetrazol-1-yl) carboxylic acids with C<sub>2</sub>–C<sub>4</sub> alkyl chains and the magnetic, structural and spectroscopic characterization of their Fe(II)–SCO complexes, as well as an initial attempt to prepare a multimetallic 3d–4f coordination polymer.

### **Results and discussion**

#### Synthesis

The spin crossover behaviour of Fe(II)–N1-alkyl-substituted tetrazole complexes has been shown to be governed by the length of the N1-alkyl substituent and the choice of weakly coordinating anion.<sup>24,31</sup> Therefore, in the current study, the length of the alkyl-skeleton was varied from  $C_2$ – $C_4$  (see Scheme 1) and  $BF_4^-$  and  $ClO_4^-$  were used as weakly coordinating tetrahedral anions of comparable volume ( $BF_4^-$  49 Å<sup>3</sup>;  $ClO_4^-$  54 Å<sup>3</sup>).

The ligands 2–4COOHTz (1–3) were prepared from the corresponding amino-acids based on the Franke-tetrazole synthesis,<sup>32</sup> in analogy to the reported synthesis of (1*H*-tetrazol-1-yl)-2-acetic acid (2).<sup>33–35</sup> Treatment of the resulting N1-alkyl-substituted tetrazoles with FeX<sub>2</sub>·6H<sub>2</sub>O (X = BF<sub>4</sub><sup>-</sup>, ClO<sub>4</sub><sup>-</sup>) in acetonitrile (MeCN) followed by subsequent washing with THF results in the desired complexes. <sup>1</sup>H NMR analysis of the bulk samples of all prepared complexes shows that only [Fe (2COOHTz)<sub>6</sub>]<sup>2+</sup> incorporated solvent molecules, for both X = BF<sub>4</sub><sup>-</sup> and ClO<sub>4</sub><sup>-</sup> amounting to 2 MeCN per Fe<sup>2+</sup> center.

#### Magnetic properties

The dependence of molar magnetic susceptibility on temperature (10 to 300 K) was investigated for all six Fe( $\pi$ )-complexes (Fig. 1a). The resulting  $\chi_{mol}T$  curves show three distinct tendencies: partial, incomplete spin crossover (5), complete spin crossover (4, 6, 7 and 9) and no spin crossover (8). All changes in  $\chi_{mol}T$  below 50 K may be attributed to zero-field splitting, expected for residual HS Fe( $\pi$ ).

 $\chi_{\rm mol}T$  decreases for 5 gradually below 170 K from 3.51 cm<sup>3</sup> K mol<sup>-1</sup> at 300 K to 2.86 cm<sup>3</sup> K mol<sup>-1</sup> at 50 K. This seems to correspond to ~25% of Fe( $\pi$ ) sites in a LS state. [Fe (2COOHTz)<sub>6</sub>](BF<sub>4</sub>)<sub>2</sub>·2 MeCN (4) shows the highest *T*<sub>1/2</sub> of the series with 227 K and decreases to a full LS-state below 180 K.



Scheme 1  $\omega$ -(1*H*-Tetrazol-1-yl) carboxylic acids used in this study, the number before the name displaying the chain-length.



View Article Online

**Dalton Transactions** 

**Fig. 1** Thermal variation of the molar magnetic susceptibility  $\chi_{mol}T$  for  $[Fe(nCOOHTz)_6]X_2$ , n = 2-4,  $X = BF_4^-$ ,  $ClO_4^-$  in the solid state between 10 K-300 K. Cooling and heating curves were found identical for all materials, so only the cooling data are displayed. (a) Overview on all materials. (b) Comparison for the solvated, partially solvated and desolvated samples of 4 and 7.

 $[Fe(4COOHTz)_6](BF_4)_2$  (6) has a  $T_{1/2}$  of 163 K, the spin-state transition taking place gradually from below 235 K to a final  $\chi_{mol}T$  of 0.98 cm<sup>3</sup> K mol<sup>-1</sup> at 50 K. This corresponds to ~72% of LS-state.

Within the ClO<sub>4</sub><sup>-</sup> compounds, [Fe(2COOHTz)<sub>6</sub>](ClO<sub>4</sub>)<sub>2</sub>·2 MeCN (7) reveals a two-step transition, featuring a kink at 170 K. Below 245 K  $\chi_{mol}T$  of 7 decreases gradually to 2.9 cm<sup>3</sup> K mol<sup>-1</sup> at 170 K, pointing to a first HS–LS transition of ~18% of the Fe(II) sites. A sharp decrease of  $\chi_{mol}T$  to a constant value of 1.2 cm<sup>3</sup> K mol<sup>-1</sup> below 115 K is attributed to further 50% of the Fe(II) sites undergoing the HS–LS transition. [Fe(3COOHTz)<sub>6</sub>] (ClO<sub>4</sub>)<sub>2</sub> (8) remains all HS-state. [Fe(4COOHTz)<sub>6</sub>](ClO<sub>4</sub>)<sub>2</sub> (9) shows by far the completest HS–LS transition with a  $T_{1/2}$  of 175 K and a final  $\chi_{mol}T$  of 0.66 cm<sup>3</sup> K mol<sup>-1</sup> at 50 K, corresponding to only ~18% of Fe(II)-centres remaining in the HS state.

The spin state transition in the  $BF_4^-$  and  $ClO_4^-$  complexes of 2COOHTz, both incorporating two molecules acetonitrile, are overly sensitive towards loss of the solvate. 4 partially loses the solvate already during drying in vacuum during the synthesis, or on shelf storage. This resulted in an intermediate with 0.7 MeCN molecules incorporated (4c, determined by <sup>1</sup>H NMR) and on subsequent drying in vacuum in a non-solvate (4b), shown in Fig. 1b. For 0.7 MeCN molecules incorporated in 4c the spin state transition is shifted to lower temperatures

#### **Dalton Transactions**

with  $T_{1/2}$  at 132 K and a final  $\chi_{mol}T$  of 2.75 cm<sup>3</sup> K mol<sup>-1</sup>, corresponding to a ~25% LS-state. The completely desolvated compound **4b** is spin crossover inactive, remaining in the HS-state for all temperatures. Similarly, the desolvated ClO<sub>4</sub><sup>-</sup>-complex **7b** shows no spin crossover anymore (Fig. 1b). This demonstrates, how crucial the incorporated solvate molecules are in this case additional to the H-bonding network.

All in all, the differing picture obtained from this comparison of the SCO-behaviour is not only caused by the weak-coordinating anion (as obvious by comparing 4 *vs.* 7, or 6 *vs.* 9), but rather attributed to a combination of different effects. Additional to the impact of the anion, as discussed in the case of 2COOHTz the 2 MeCN solvate makes the game. Furthermore, there is the variation in the length of the alkyl-spacer, which for 4COOHTz seems to be a good fit between establishing cooperativity and preventing too much motional freedom in form of a shock-absorber effect,<sup>24,31</sup> which was evidenced in the past for increasing chain-lengths. Missing the structural characterization of the 3COOHTz-compounds, the differing SCO-behaviour is probably related to the odd number of C-atoms, which already for unsubstituted alkyl-tetrazoles were found challenging in the past.<sup>24,31</sup>

#### Crystallographic analysis

Single crystals suitable for determination of the molecular structure could be grown for **4**, **6**, 7 and **9** by vapour-diffusion of diethyl ether ( $Et_2O$ ) into a concentrated MeCN-solution of the corresponding compounds. No suitable crystals of **5** and **8** could be obtained by similar means. Single crystal diffraction data were collected at 100 and 200 K for **4**, **6**, 7 and **9** in order to determine both low- and high-spin molecular structures. In addition variable temperature P-XRD patterns were acquired for **4** and 7 from 100–300 K.

[Fe(2COOHTz)<sub>6</sub>](BF<sub>4</sub>)<sub>2</sub>·2 MeCN (4). 4 crystallizes as MeCN solvate in the trigonal  $R\bar{3}$  space group and this symmetry is retained both at 100 K (low-spin, see Fig. 2) and 200 K (highspin). The iron-centre is octahedrally surrounded by six 2COOHTz-ligands coordinating via the exo N4-nitrogen. The Fe-N bond lengths for all six ligands are equal (1.987 Å), typical for a Fe(II) LS-state. At 200 K the distance increases ~9% to 2.176 Å, characteristic for the Fe( $\pi$ ) HS-state (see Table S1<sup>†</sup>).<sup>1</sup> The packing in the crystal is governed – as predicted for a COOH-group - by an H-bond network composed of infinite C(5) type chains between the COOH-groups. Viewed along the *b* axis, the  $[Fe(2COOHTz)_6]^{2+}$ -cations stack in layers, with three of the ligands pointing upwards and three downwards (see Fig. 2) to the adjacent layer. This arrangement is further stabilised by the above-mentioned H-bonds (see Fig. 3). The Fe-atoms are located on the Wyckoff-position 3a, alternating with an acetonitrile, an apex-up and an apex-down  $BF_4$  group. The  $BF_4$  groups interact strongly with the  $[Fe(2COOHTz)_6]^{2+}$ -cations through C-H···F-bonds with both the tetrazolic CH and CH<sub>2</sub>-group (see Fig. 4) in both the highand low-spin states.

[Fe(2COOHTz)6](ClO4)2.2 MeCN (7). 7 crystallizes isotypically to 4 with slightly larger cell parameters (see Table 1) as



Fig. 2 Molecular structure of 4 at 100 K in the LS-state showing H-bond contacts at the carboxylic acid groups; H-atoms omitted for clarity.



Fig. 3 H-Bonding network in 4 at 100 K, seen along the *b*-axis.



Fig. 4  $H \cdots F$  bonds between the tetrazolic CH and the CH<sub>2</sub>-groups in 4 at 100 K.

an MeCN solvate in the trigonal  $R\bar{3}$  space group. In 7 the Fe–N distances (2.004 Å, 100 K, LS-state; 2.184 Å, 200 K, HS-state, see Table S2†) are slightly longer than in 4, but still compar able and characteristic for Fe( $\pi$ ) in the corresponding spin-state. Also, in 7 both at 100 K and 200 K the ClO<sub>4</sub><sup>-</sup>-anion is not disordered and stabilized through interaction with the [Fe (2COOHTz)<sub>6</sub>]<sup>2+</sup>-cation (see Fig. 5).

Variable temperature P-XRD of bulk (4) and (7). Magnetic measurements of bulk 4 and 7 show an incomplete spin transition, whereas the Fe–N bond distances obtained from the single crystal data suggest a largely complete transition. Therefore, variable temperature P-XRD was used to follow any structural changes of the bulk material during the SCO (see Fig. 6). For both compounds, the calculated diffractograms obtained from the HS and LS molecular structures are in reasonable agreement with the powder-patterns of the bulk material. For both 4 (Fig. 6a) and 7 (Fig. 6b) the temperature dependent PXRD measurements show more pronounced changes in the powder pattern (dotted lines as a guide) than to those just expectable from thermal contraction. On this basis it seems clear that both bulk and single crystalline samples are the same compound.

 $[Fe(4COOHTz)_6](BF_4)_2 \cdot Et_2O$  (6). 6 crystallizes as an  $Et_2O$ solvate, as a result of vapour-diffusion crystallization, in the triclinic P1 space group. The structure was determined at 100 K (LS-state) and 200 K (HS-state). The asymmetric unit (see Fig. 7) includes the Fe-centre with three 4COOHTz ligands, one crystallographic independent BF4-anion, as well as half of the Et<sub>2</sub>O-solvate. The Fe-N distances of the three coordinated ligands are slightly different (1.961 Å (N4), 1.979 Å (N4a) and 1.989 Å (N4b), see Table S3<sup>†</sup>). In the HS-state at 200 K the Fe-N distances increase to 2.185 Å (N4), 2.182 Å (N4a) and 2.185 Å (N4b). As for 4 and 7, the supramolecular structure is governed by an H-bond network with the COOHgroups forming a three-dimensional network of  $R_2^2$  (8) type ring motifs joining layers of [Fe(4COOHTz)<sub>6</sub>]<sup>2+</sup>-cations (see Fig. 8). The voids between the ligand alkyl-chains are occupied by  $Et_2O$ , which also interacts with the COOH-groups via a

single H-bond. Comparing the supramolecular arrangement of 4/7 to **6** it becomes evident, that in this case only due to the longer alkyl-chains Et<sub>2</sub>O could be incorporated as solvate molecule. In the former case the voids between the cationic Fe-layers would not have been large enough to fit the Et<sub>2</sub>O-molecules. Whereas at 100 K the BF<sub>4</sub><sup>-</sup>-anions are stabilized by the H…F bonds similar to 4 preventing a potential disorder, at 200 K in the HS-state those interactions are no longer strong enough, resulting in an apex up-down disorder of the BF<sub>4</sub><sup>-</sup>-groups.

Structure of [Fe(4COOHTz)<sub>6</sub>](ClO<sub>4</sub>)<sub>2</sub>·Et<sub>2</sub>O (9). 9 crystallizes as Et<sub>2</sub>O solvate like 6 in the triclinic  $P\bar{1}$  space group but has two crystallographic independent Fe-centres and two ClO<sub>4</sub><sup>-</sup>-anions in the asymmetric unit. Both Fe1 and Fe2 are surrounded by each three independent 4COOHTz-ligands, which do not interact with each other via H-bonds. The solvate molecule is located in the cavity formed by the ligands, stabilized by an H-bond (see Fig. 9). At 100 K, all Fe–N bond lengths (1.984 Å Fe1-N4, 1.987 Å Fe1-N4a, 1.982 Å Fe1-N4b, 1.987 Å Fe2N4c, 1.987 Å Fe2–N4d, 1.984 Å Fe2–N4e) are characteristic for Fe(π) in its low-spin state. In contrast, at 200 K the bonds extend to typical HS distances (2.181 Å Fe1–N4, 2.178 Å Fe1–N4a, 2.173 Å Fe1-N4b, 2.174 Å Fe2N4c, 2.185 Å Fe2-N4d, 2.178 Å Fe2-N4e, see Table S4<sup>†</sup>). In **9** the  $[Fe(4COOHTz)_6]^{2+}$ -cations form layers connected through the H-bond network to each other and the ClO<sub>4</sub><sup>-</sup>-anions.

The relevant crystallographic parameters for all Fe-containing structures are given in Table 1.

#### Spectroscopic characterization

MIR and FIR spectroscopy. For homoleptic Fe(II)-tetrazolecomplexes IR-spectroscopy allows for confirmation of successful SCO-complex formation, as well as gaining insight into the spin state present, since the tetrazolic CH-vibration is sensitive to the N4 coordination environment.36 In Fig. 10, the IR spectra of 2COOHTz (1) and its  $ClO_4^{-}$ -complex (7) in the HSstate (red) and LS-state (blue) are compared. At 3156 cm<sup>-1</sup> the free ligand shows its tetrazolic CH-vibration, which upon successful coordination is split and shifted to 3159 cm<sup>-1</sup> and 3152 cm<sup>-1</sup>. On cooling to the LS-state, at 100 K the CHvibration is shifted to 3164 cm<sup>-1</sup> and 3159 cm<sup>-1</sup>, respectively. Around 1600 cm<sup>-1</sup> the  $\nu_{\rm N2=N3}$  stretching vibration of the tetrazole after coordination is similarly affected by the spin-state transition, shifting from 1612 cm<sup>-1</sup> to 1616 cm<sup>-1</sup> in the LSstate. The change of the carbonyl-vibration at 1731  $\text{cm}^{-1}$  on coordination and afterwards during SCO may be attributed to interaction within the H-bond network, transferring the volume work of the spin-state transition. The broad absorption around 1030 cm<sup>-1</sup> corresponds to the B-F stretching vibrations of the BF<sub>4</sub><sup>-</sup>-anion. The CN-absorption of the MeCN solvate in 7 is very weak, and is only observable in the LS-spectrum of 7 in Fig. 10 around  $2250 \text{ cm}^{-1}$ .

In the FIR-region of the spectra, both spin-states show the characteristic displacement of the Fe-centre towards the centroid of the trigonal faces of the N6-coordination octahedron.

#### **Dalton Transactions**

	4·MeCN, HS	4·MeCN, LS	7∙MeCN, HS	7∙MeCN, LS
Formula	$C_{22}H_{30}B_2F_8FeN_{26}O_{12}$	C22H30B2F8FeN26O12	C22H30Cl2FeN26O20	C22H30Cl2FeN26O20
Weight [g mol <sup>-1</sup> ]	1080.19	1080.19	1105.47	1105.47
T [K]	200	100	200	100
Colour	White	Pink	White	Pink
Shape	Platelet	Platelet	Platelet	Platelet
Crystal system	Trigonal	Trigonal	Trigonal	Trigonal
Space group	RĪ	RĪ	RĪ	RĪ
<i>a</i> [Å]	10.6588(7)	10.4735(5)	10.747(3)	10.587(2)
b Å	10.6588(7)	10.4735(5)	10.747(3)	10.587(2)
<i>c</i> [Å]	34.838(3)	34.480(2)	34.689(9)	34.314(7)
α <sup>[</sup> o]	90	90	90	90
β <sup>[o]</sup>	90	90	90	90
γ[0]	120	120	120	120
V[Å <sup>3</sup> ]	3427.7(5)	3275.6(4)	3470(2)	3330.8(14)
Z	3	3	3	3
$\rho_{\rm rate}  \left[ g  {\rm cm}^{-3} \right]$	1.570	1.643	1.587	1.653
$\mu \left[ \text{mm}^{-1} \right]$	0.445	0.466	0 543	0.565
Measured refl's	22 250	21 166	27.060	26.274
Indep't refl's	1572	1405	1025	1950
Pefl's $I > 2\sigma(I)$	1421	1274	1335	1855
Refit S $I \ge 20(I)$	1421	1374	1389	1444
R <sub>int</sub>	0.0308	0.0309	0.0952	0.0798
GOOF	1.068	1.072	1.0/1	1.115
WK <sub>2</sub>	0.0804	0.06/3	0.1449	0.1329
$R_1$	0.0318	0.02/1	0.0569	0.0550
CCDC†	2024140	2024142	2024146	2024147
	$6 \cdot \text{Et}_2 \text{O}$ , HS	$6 \cdot \text{Et}_2 \text{O}$ , LS	$9 \cdot Et_2O$ , HS	$9 \cdot \text{Et}_2 \text{O}$ , LS
Formula	C34H58B2F8FeN24O13	C34H58B2F8FeN24O13	C <sub>34</sub> H <sub>58</sub> Cl <sub>2</sub> FeN <sub>24</sub> O <sub>21</sub>	C <sub>34</sub> H <sub>58</sub> Cl <sub>2</sub> FeN <sub>24</sub> O <sub>21</sub>
Weight [g mol <sup>-1</sup> ]	1240.45	1240.45	1265.79	1265.79
T[K]	200	100	200	100
Colour	White	Pink	White	Pink
Shape	Platelet	Platelet	Platelet	Platelet
Crystal system	Triclinic	Triclinic	Triclinic	Triclinic
Space group	$P\bar{1}$	$P\bar{1}$	$P\bar{1}$	$P\bar{1}$
a[Å]	10.630(3)	10.530(3)	10.6262(13)	10.6120(12)
b [Å]	10.971(3)	10.576(3)	11.0192(13)	10.6186(12)
	14.235(4)	13.836(4)	27.044(3)	26.596(3)
a [0]	74 796(6)	75 400(8)	96.080(4)	95 237(3)
β[0]	71 624(6)	73 122(7)	93 160(4)	95.276(3)
<i>P</i> [] <i>v</i> [0]	63 563(6)	64.974(7)	116 571(3)	114 758(3)
7 L J V [Å <sup>3</sup> ]	1206 1(6)	1201 0(7)	2707 5(6)	2691 0(5)
v [A ]	1390.1(0)	1321.2(7)	2797.5(0)	2001.9(3)
$a [a am^{-3}]$	1 510	1 665	1 502	1 567
$\rho_{\text{calc.}}[g \text{ cm}]$	1.312	1.005	1.505	1.307
$\mu$ [IIIIII ]	0.377	0.396	0.459	0.479
measured refl's.	32 025	28 095	01 130	53/54
indepit reflis	6916	6513	13 866	13 356
Refl's $l \ge 2\sigma(l)$	2942	2/33	640/	6404
R <sub>int</sub>	0.1497	0.2107	0.0757	0.0644
GooF	1.014	1.002	1.051	1.010
$wR_2$	0.3257	0.1728	0.2197	0.1016
$R_1$	0.1327	0.0910	0.0874	0.0596
-				

Those vibrations are nearly entirely decoupled from other vibrational modes and therefore indicative of spin state. For the HS-state, this vibration is found at 226 cm<sup>-1</sup>, for the LS-state the characteristic vibrations are located at 470 and 443 cm<sup>-1</sup> (see Fig. 11).

UV-VIS/NIR spectroscopy. Electronic transitions are also affected by the spin state transition. For homoleptic Fe-tetrazole SCO complexes the HS-state absorbs around 850 nm in the near-infrared, causing a broad and often weak absorption in the ligand field spectrum. This  ${}^{5}T_{2} \rightarrow {}^{5}E$  transition weakens with cooling and spin-state transition, giving rise to two

absorptions in the visible region of the spectrum at 545 nm and 386 nm, responsible for the pink colour of the LS-state. These absorption bands correspond to the  ${}^{1}A_{1} \rightarrow {}^{1}T_{1}$  transition, as well as the  ${}^{1}A_{1} \rightarrow {}^{1}T_{2}$  transition. From the maxima observed in the HS and the LS state, a ligand-field strength 10 Dq can be estimated.<sup>37</sup> In the case of 7, measured in the solid state, a value of 1.65 Dq is obtained (Fig. 12).

#### Towards a mixed-metallic 3d-4f coordination polymer

The functionalization of alkyl-tetrazoles with a COOH-group introduces an H-bond donor-acceptor system thereby enhan-



Fig. 5 Hydrogen-bonds between the tetrazolic CH and the  $ClO_4^-$ -anion, as well as short interactions with the  $CH_2$ -group in 7 at 100 K.



Fig. 6 Temperature-dependent P-XRD data for (a)  $[Fe(2COOHTz)_6]$   $(BF_4)_2 \cdot 2MeCN$  (4) and (b)  $[Fe(2COOHTz)_6](CIO_4)_2 \cdot 2MeCN$  (7).

cing supramolecular cooperativity of SCO. To take advantage of the coordination ability of the COOH-group and the bifunctional ligand, we attempted to prepare a multi-metallic 3d–4f coordination polymer. Coronado *et al.* have success-



Fig. 7 Asymmetric unit of [Fe(4COOHTz)<sub>6</sub>](BF<sub>4</sub>)<sub>2</sub>·Et<sub>2</sub>O (6) at 100 K.



Fig. 8 Supramolecular H-bonding network in 6 at 100 K, seen along the a-axis.

fully applied such an approach to the formation of a mixed Fe(III)–Fe(II) SCO compound using a COOH-functionalized ligand.<sup>38</sup>

By reacting 3COOHTz (2) with  $Fe(ClO_4)_2 \cdot 6H_2O$  and GdCl<sub>3</sub>·6H<sub>2</sub>O in aqueous MeOH and subsequent evaporation a slightly yellow oil was obtained, which did not be solidify. After a few weeks at 4 °C a few colourless crystals had formed, allowing for determination of the structure by X-ray diffraction. Instead of a mixed-metallic coordination polymer, [Gd  $(3COOTz)_2(H_2O)_3$  Cl (10) had crystallized in the monoclinic C2/ c space group. During the reaction/crystallization process the COOH-groups were deprotonated and the resulting carboxylate groups act as chelating, bridging ligands to the gadoliniumatoms in a  $\mu_2$ - $\eta^2$ : $\eta^1$  fashion (see Fig. 13). Each gadolinium atom is coordinated nine-fold by oxygen-donors as the centre of a monocapped square antiprismatic coordination polyhedron. Due to the  $\mu_2$ - $\eta^2$ : $\eta^1$  coordination of the carboxylateligands one-dimensional chains are formed, in which the Gd-atom is coordinated by four 3COOTz-ligands: two of them



Fig. 9 Asymmetric unit of [Fe(4COOHTz)<sub>6</sub>](ClO<sub>4</sub>)<sub>2</sub>·Et<sub>2</sub>O (9) at 100 K.



Fig. 10 Comparison of MIR spectra for 2COOHTz (1),  $[Fe(2COOHTz)_6]$  (ClO<sub>4</sub>)<sub>2</sub>·2 MeCN (7) in the HS-state (red) and LS-state (blue).



Fig. 11 Comparison of FIR spectra for  $[Fe(2COOHTz)_6](ClO_4)_2$ ·2MeCN (7) in the HS-state (red) and LS-state (blue).



Fig. 12 Temperature-dependent UV-VIS/NIR spectra of [Fe (2COOHTz)<sub>6</sub>](ClO<sub>4</sub>)<sub>2</sub>·2MeCN (7). Inset: Increase of the absorption corresponding to the  $^1A_1 \rightarrow \,^1T_1$  transition on cooling and increasing number of LS Fe-centres.



Fig. 13  $\mu_2$ - $\eta^2$ : $\eta^1$  coordination pattern of the carboxylate-ligands in [Gd (3COOTz)<sub>2</sub>(H<sub>2</sub>O)<sub>3</sub>]Cl (10).

coordinate as bidentate, and one 3COOTz-ligand of each adjacent Gd-centre on both sides coordinates in an  $\eta^1$ -fashion (see Fig. 14). H<sub>2</sub>O molecules occupy the remaining coordination sites (3). The  $\eta^1$ -interaction of the carboxylate-ligands with the adjacent Gd-atoms is remarkable, as the Gd–O-bond is the shortest of all three: the  $\eta^1$  Gd–O interaction is 2.350 Å (Gd–O1), whereas the  $\eta^2$  Gd–O bonds are with 2.432 Å (Gd–O2) and 2.685 Å (Gd–O1) 3.48% and 14.3% longer. The tetrazoles establish channels parallel to the gadolinium-oxo chains. Their tetrazolic CH groups are directed to the inner of the voids, the N2 and N3 atoms of the ring interacting with the H<sub>2</sub>O ligands of the nearby gadolinium-oxo chain. In the voids the Cl<sup>-</sup>-anions are stabilized by a zig-zag H-bond structure, originating from the H<sub>2</sub>O-ligands at the Gd-centre (see Fig. 15). The crystallographic parameters for **10** are given in Table 2.



**Fig. 14** One-dimensional chains in  $[Gd(3COOT_z)_2(H_2O)_3]Cl$  (**10**) with  $Cl^-$ -atoms parallel to the *c*-axis, seen along the *a*-axis.



Fig. 15 Cl<sup>-</sup>-Atoms located in the voids formed by the tetrazole rings stabilized by H-bonds to the  $H_2O$  ligands in the molecular structure of 10, view along *c*-axis.

Table 2 Crystallographic parameters for 10

	10	
Formula	C <sub>8</sub> H <sub>16</sub> ClGdN <sub>8</sub> O <sub>7</sub>	
Weight $[g mol^{-1}]$	528.96	
T [K]	100	
Colour	Clear colourless	
Shape	Plate	
Crystal system	Monoclinic	
Space group	C2/c	
a [Å]	18.0057(4)	
b [Å]	11.2323(6)	
c [Å]	8.0169(9)	
α [°]	90	
$\beta$ [°]	91.838(3)	
γ [°]	90	
$V[Å^3]$	1620.5(2)	
z	4	
$r_{\rm calc.} [\rm g \ \rm cm^{-3}]$	2.1674	
$m [\mathrm{mm}^{-1}]$	4.311	
Measured refl's	9983	
Indep't refl's	2927	
Refl's $I \ge 2s(I)$	2505	
R <sub>int</sub>	0.0618	
GooF	1.67	
wR <sub>2</sub>	0.0367	
R <sub>1</sub>	0.0331	
CCDC†	2024832	



Fig. 16 Comparison of MIR spectra of 3COOHTz (2) and [Gd  $(3COOTz)_2(H_2O)_3lCl$  (10).

Fig. 16 shows a comparison between the MIR spectrum of 2 and a single crystal of  $[Gd(3COOTz)_2(H_2O)_3]Cl$ . Whereas the tetrazolic CH-band is only slightly shifted after the coordination to the  $Gd^{3+}$ , the deprotonated and coordinated carbonyl-vibration is shifted by 20 cm<sup>-1</sup>. The three coordinated H<sub>2</sub>O-ligands appear in the MIR, resulting the OH-stretching mode at 3346 cm<sup>-1</sup> and the OH-scissoring vibration at 1672 cm<sup>-1</sup>.

## Experimental

#### Materials and methods

All operations involving Fe(n) were carried out under inert gas atmosphere (argon 5.0). The glassware used was oven dried at 120 °C before use for at least 2 hours. All solvents for the complexation reactions were dried before use and stored over molecular sieve 3 Å under argon.<sup>39</sup> Unless otherwise stated, all starting materials were commercially obtained and used without further purification. All NMR spectra were recorded in dry deuterated solvents on a Bruker Avance UltraShield 400 MHz. Chemical shifts are reported in ppm; <sup>1</sup>H and <sup>13</sup>C shifts are referenced against the residual solvent resonance. For the measurement of MIR and FIR spectra, a PerkinElmer Spectrum 400, fitted with a coolable/heatable PIKE Gladi ATR unit was used within the range of 4000-100 cm<sup>-1</sup>. Solid state UV/Vis/NIR spectra were recorded with a PerkinElmer Lambda 900 spectrophotometer between 300 and 1600 nm in diffuse reflectance against BaSO<sub>4</sub>. A Harrick coolable/heatable powder sample holder in "Praying Mantis" configuration was used. Melting points were determined by differential scanning calorimetry, using a Netzsch STA 449 C Jupiter® with heating rates of 2 K min<sup>-1</sup>, see Fig. S1–S3.† The magnetic moment of the Fe(II) complexes was measured using a Physical Property Measurement System (PPMS®) by Quantum Design. The

experimental setup consisted of a vibrating sample magnetometer attachment (VSM), bearing a brass sample holder with a quartz-glass powder container. The magnetic moment was determined in an external field of 1 T in the range of 10 K to 300 K, measuring all 5 K with a previous thermal stabilization of 5 minutes. Variable temperature mid-range  $(4000-450 \text{ cm}^{-1})$ infrared spectra were recorded by the ATR technique on a PerkinElmer Spectrum 400, fitted with a coolable/heatable PIKE Gladi ATR Unit.<sup>25</sup> Single crystals were attached to a glass fiber by using perfluorinated oil and were mounted on a Bruker KAPPA APEX II diffractometer equipped with a CCD detector with Mo  $K_{\alpha}$  radiation (Incoatec Microfocus Source IµS: 30 W, multilayer mirror,  $\lambda = 0.71073$  Å). For all measurements data were reduced to intensity values by using SAINT Plus,<sup>40</sup> and an absorption correction was applied by using the multi scan method implemented by SADABS.<sup>40</sup> For the iron(II) complexes, protons were placed at calculated positions and refined as riding on the parent C atoms. All non-H atoms were refined with anisotropic displacement parameters. For 2024832 (10), a Bruker KAPPA APEX II diffractometer equipped with a CCD detector was used and data were collected at 100 K. The powder X-ray diffraction measurements were carried out on a PANalytical X'Pert Pro diffractometer in Bragg-Brentano geometry using Cu  $K_{\alpha 1,2}$  radiation filtered with a BBHD mirror and an X'Celerator linear detector. For in situ experiments below ambient temperature an Oxford PheniX Cryochamber from Oxford Cryosystems was used. The powder sample were mounted on a copper sample holder on top of a background-free silicon support. The sample chamber was evacuated, and all measurements were carried out under vacuum. The actual sample temperature is directly monitored by a thermocouple on the sample holder. The diffractograms were evaluated using the PANalytical program suite HighScorePlus, correcting for the background.

#### Synthesis of ligands

Tetrazoles and their derivatives – especially in combination with perchlorates – are potentially shock sensitive or explosive compounds and should, therefore, be handled with great care and with the appropriate safety precautions!

**2-(1H-Tetrazol-1-yl)acetic acid (2COOHTz) (1).** 25 g (333 mmol, 1 eq.) glycine and 32.4 g (499.6 mmol, 1.5 eq.) NaN<sub>3</sub> were suspended in 97 ml (516.2 mmol, 1.55 eq.) triethylorthoformate and 250 ml acetic acid. The suspension was heated for 18 h at 95 °C, filtrated and evaporated to dryness. After extracting the residue with CH<sub>2</sub>Cl<sub>2</sub>, the solvent was removed, and the resulting yellow oil refluxed for 6 h in concentrated hydrochloric acid. After removal of all volatiles the oily residue was adsorbed on SiO<sub>2</sub> and was further purified by column chromatography (MeCN : CH<sub>2</sub>Cl<sub>2</sub> = 3 : 2). After evaporation, **1** was obtained as beige crystalline material. Yield: 7.9 g, 18.5%; MP: 130 °C;  $\nu_{CH(Tz)}$  3156,  $\nu_{CO}$  1731/1712 cm<sup>-1</sup>; <sup>1</sup>H NMR (400 MHz, DMSO-d<sub>6</sub>)  $\delta$  = 13.43 (s, 1H, COOH), 9.38 (s, 1H, Tz), 5.43 (s, 2H, CH<sub>2</sub>) ppm; <sup>13</sup>C{<sup>1</sup>H} NMR (101 MHz, DMSO-d<sub>6</sub>)  $\delta$  = 167.94 (COOH), 144.99 (Tz), 48.52 (CH<sub>2</sub>) ppm.

3-(1H-Tetrazol-1-yl)propanoic acid (3COOHTz) (2). 25 g (280.6 mmol, 1 eq.) β-alanine and 27.4 g (420.9 mmol, 1.5 eq.) NaN3 were suspended in 81.7 ml (434.9 mmol, 1.55 eq.) triethylorthoformate and 250 ml acetic acid. The suspension was heated for 18 h at 95 °C, filtrated and evaporated to dryness. After extracting the residue with CH<sub>2</sub>Cl<sub>2</sub>, the solvent was removed, and the resulting yellow oil refluxed for 6 h in concentrated hydrochloric acid. After removal of all volatiles the oily residue was adsorbed on SiO<sub>2</sub> and was further purified by column chromatography (MeCN:  $CH_2Cl_2 = 3:2$ ). After evaporation, 3COOHTz was obtained as off-white solid. Yield: 9.1 g, 22.8%; MP: 125 °C;  $\nu_{CH(Tz)}$  3134,  $\nu_{CO}$  1709 cm<sup>-1</sup>; <sup>1</sup>H NMR (400 MHz, DMSO-d<sub>6</sub>)  $\delta$  = 12.34 (s, 1H, COOH), 9.37 (s, 1H, Tz), 4.64 (t, J = 6.6 Hz, 2H, CH<sub>2</sub>), 2.98 (t, J = 6.7 Hz, 2H, CH<sub>2</sub>) ppm; <sup>13</sup>C{<sup>1</sup>H} NMR (101 MHz, DMSO-d<sub>6</sub>)  $\delta$  = 171.72 (COOH), 144.30 (Tz), 43.74 (CH<sub>2</sub>), 33.49 (CH<sub>2</sub>).

4-(1H-Tetrazol-1-yl)butanoic acid (4COOHTz) (3). 25 g (242.4 mmol, 1 eq.) 4-aminobutanoic acid and 23.6 g (363.7 mmol, 1.5 eq.) NaN<sub>3</sub> were suspended in 70.6 ml (375.8 mmol, 1.55 eq.) triethylorthoformate and 250 ml acetic acid. The suspension was heated for 18 h at 95 °C, filtrated and evaporated to dryness. After extracting the residue with CH<sub>2</sub>Cl<sub>2</sub>, the solvent was removed, and the resulting yellow oil refluxed for 6 h in concentrated hydrochloric acid. After removal of all volatiles the oily residue crystallized in the freezer overnight. The yellow solid was adsorbed on SiO2 and further purified by column chromatography (MeCN :  $CH_2Cl_2 =$ 3:2) After evaporation, 4COOHTz was obtained as off-white solid. Yield: 11.24 g, 29.7%; MP: 86 °C;  $\nu_{CH(Tz)}$  3114,  $\nu_{CO}$ 1714 cm<sup>-1</sup>; <sup>1</sup>H NMR (400 MHz, DMSO-d<sub>6</sub>)  $\delta$  = 12.21 (s, 1H, COOH), 9.41 (s, 1H, Tz), 4.48 (t, J = 7.1, 2H, CH<sub>2</sub>), 2.27 (t, J = 7.3, 2H, CH<sub>2</sub>), 2.06 (p, J = 7.2, 2H, CH<sub>2</sub>) ppm; <sup>13</sup>C{<sup>1</sup>H} NMR (101 MHz, DMSO-d<sub>6</sub>)  $\delta$  = 173.52 (COOH), 143.99 (Tz), 46.91 (CH<sub>2</sub>), 30.37 (CH<sub>2</sub>), 24.72 (CH<sub>2</sub>).

#### Synthesis of Fe(II)-complexes

[Fe(2COOHTz)<sub>6</sub>][BF<sub>4</sub>]<sub>2</sub>·2MeCN (4). Fe(BF<sub>4</sub>)2·6H<sub>2</sub>O (100 mg, 296.3 µmol, 1 eq.) and 1 (231.5 mg, 1.81 mmol, 6.1 eq.) were dissolved in 10 ml MeCN and stirred at 50 °C for 18 h. After evaporation of the solvent, the residual oil was triturated in 10 ml THF for 1 h to remove the excess of ligand. The precipitated material was separated, washed with further THF (10 ml) and dried in a stream of N<sub>2</sub>. Yield: 98.8 mg, 33.4%;  $\nu_{CH(Tz)}$  3138,  $\nu_{CO}$  1744 cm<sup>-1</sup>.

[Fe(3COOHTz)<sub>6</sub>][BF<sub>4</sub>]<sub>2</sub> (5). Same procedure as for 4. 5 was isolated as off-white, sticky solid. Yield: 75.7 mg, 23.6%;  $\nu_{CH(Tz)}$  3135,  $\nu_{CO}$  1714 cm<sup>-1</sup>.

[Fe(4COOHTz)6][BF4]2 (6). Same procedure as for 4. 6 was isolated as white solid. Yield: 122 mg, 35.3%;  $\nu_{CH(Tz)}$  3144,  $\nu_{CO}$  1733 cm<sup>-1</sup>.

[Fe(2COOHTz)<sub>6</sub>][ClO<sub>4</sub>]<sub>2</sub>·2MeCN (7). Fe(ClO<sub>4</sub>)<sub>2</sub>·6H<sub>2</sub>O (100 mg, 275.6  $\mu$ mol, 1 eq.) was dissolved under Ar together with a tiny amount of ascorbic acid in 2 ml MeCN and added by filtration through a syringe filter (PTFE, 0.45  $\mu$ m) to a degassed solution of 1 (215.4 mg, 1.68 mmol, 6.1 eq.) in 8 ml MeCN. The resulting mixture was stirred at 50 °C for 18 h. After evaporation of

**Dalton Transactions** 

the solvent, the residual oil was triturated in 10 ml THF for 1 h to remove the excess of ligand. The precipitated material was separated, washed with further THF (10 ml) and dried in a stream of N<sub>2</sub>. Yield: 51.5 mg, 18.2%;  $\nu_{CH(Tz)}$  3159/3152,  $\nu_{CO}$  1733/1715 cm<sup>-1</sup>.

[Fe(3COOHTz)<sub>6</sub>][ClO<sub>4</sub>]<sub>2</sub> (8). Same procedure as for 7. 8 was isolated as white sticky solid. Yield: 127.1 mg, 41.6%;  $\nu_{CH(Tz)}$  3137,  $\nu_{CO}$  1740/1715 cm<sup>-1</sup>.

[Fe(4COOHTz)<sub>6</sub>][ClO<sub>4</sub>]<sub>2</sub> (9). Same procedure as for 7. 9 was isolated as white solid. Yield: 118.9 mg, 36.2%;  $\nu_{CH(Tz)}$  3134,  $\nu_{CO}$  1728 cm<sup>-1</sup>.

[Gd(3COOTz)<sub>2</sub>(H<sub>2</sub>O)<sub>3</sub>]Cl (10). Fe(ClO<sub>4</sub>)<sub>2</sub>·6H<sub>2</sub>O (42.55 mg, 117.27  $\mu$ mol, 1 eq.) was dissolved under Ar together with a tiny amount of ascorbic acid in 2 ml MeOH and added by filtration through a syringe filter (PTFE, 0.45  $\mu$ m) to a degassed solution of 2 (100 mg, 703.64 mmol, 6 eq.) in 8 ml of a 1:1 mixture MeOH:H<sub>2</sub>O. The resulting mixture was stirred at 50 °C for 10 minutes, then 43.6 mg (117.27  $\mu$ mol, 1 eq.) GdCl<sub>3</sub>·6H<sub>2</sub>O was added. The mixture was further stirred for 6 h at 50 °C. After evaporation of the solvent, the residual slightly yellow oil was kept at 50 °C for 6 h under vacuum and afterwards left for 3 weeks at 4 °C. A few tiny colourless crystals formed.

# Conclusions

Fe(π)-SCO complexes with bifunctional ω-(1H-tetrazol-1-yl) carboxylic acids allow formation of supramolecular hydrogen bonding networks that enhance the cooperativity of the material. Furthermore, the COOH-group can act as anchorgroup for deposition on oxidic surfaces, or for formation of multi-metallic coordination polymers. Three ligands bearing a COOH-group with alkyl-chain lengths of C2-C4 were synthesized and coordinated to Fe(II) with  $BF_4^-$  and  $ClO_4^-$  as weak-coordinating anions. Complete and sharp spin state transitions were observed in the cases of 4, 6, 7 and 9. An initial attempt to prepare a 3d-4f mixed-metallic coordination polymer with Gd<sup>3+</sup> resulted in the unexpected formation of one-dimensional Gd-oxo chains with the 3COOHTz-ligand having been deprotonated and coordinating to the Gd<sup>3+</sup>-atoms in a chelating-bridging  $\mu_2$ - $\eta^2$ : $\eta^1$  fashion. Future work will focus on a more suitable synthetic approach to realise a coordination on both ligand functional groups.

# Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

We acknowledge financial support of the Austrian Science Fund (FWF Der Wissenschaftsfond) project P 31076-N28. The X-Ray center (XRC) of the Vienna University of Technology provided access to the powder X-Ray diffractometer.

### References

- 1 P. Gütlich, H. A. Goodwin, Topics in Current Chemistry, *Spin Crossover in Transition Metal Compounds I–III*, Springer Berlin Heidelberg, 2004.
- 2 P. Gütlich, *Eur. J. Inorg. Chem.*, 2013, 581–591, DOI: 10.1002/ejic.201300092.
- 3 H. J. Shepherd, C. Bartual-Murgui, G. Molnar, J. A. Real, M. C. Munoz, L. Salmon and A. Bousseksou, *New J. Chem.*, 2011, 35, 1205–1210.
- 4 S. Decurtins, P. Gutlich, C. P. Kohler, H. Spiering and A. Hauser, *Chem. Phys. Lett.*, 1984, **105**, 1–4.
- 5 A. C. Aragonès, D. Aravena, J. I. Cerdá, Z. Acís-Castillo, H. Li, J. A. Real, F. Sanz, J. Hihath, E. Ruiz and I. Díez-Pérez, *Nano Lett.*, 2016, **16**, 218–226.
- 6 C. Lefter, V. Davesne, L. Salmon, G. Molnár, P. Demont, A. Rotaru and A. Bousseksou, *Magnetochemistry*, 2016, 2, 18.
- 7 C. Lefter, S. Rat, J. S. Costa, M. D. Manrique-Juárez, C. M. Quintero, L. Salmon, I. Séguy, T. Leichle, L. Nicu, P. Demont, A. Rotaru, G. Molnár and A. Bousseksou, *Adv. Mater.*, 2016, 28, 7508–7514.
- 8 C. Lefter, R. Tan, J. Dugay, S. Tricard, G. Molnár, L. Salmon, J. Carrey, W. Nicolazzi, A. Rotaru and A. Bousseksou, *Chem. Phys. Lett.*, 2016, 644, 138–141.
- 9 L. Cambi and A. Cagnasso, Atti R. Accad. Naz. Lincei, Mem. Cl. Sci. Fis., Mat. Nat., 1931, 13, 809.
- 10 L. Cambi and L. Szegö, Ber. Dtsch. Chem. Ges. (A and B), 1933, 66, 656–661.
- 11 L. Cambi and L. Szegö, Ber. Dtsch. Chem. Ges. (A and B), 1931, 64, 2591–2598.
- 12 M. Ohba, K. Yoneda and S. Kitagawa, *CrystEngComm*, 2010, 12, 159–165.
- 13 C. J. Kepert, *Chem. Commun.*, 2006, 695, DOI: 10.1039/ b515713g.
- 14 A. Holovchenko, J. Dugay, M. Giménez-Marqués, R. Torres-Cavanillas, E. Coronado and H. S. J. van der Zant, *Adv. Mater.*, 2016, 28, 7228–7233.
- 15 J. Dugay, M. Gimenez-Marques, T. Kozlova, H. W. Zandbergen, E. Coronado and H. S. J. van der Zant, *Adv. Mater.*, 2015, 27, 1288–1293.
- 16 H. J. Shepherd, I. Y. A. Gural'skiy, C. M. Quintero, S. Tricard, L. Salmon, G. Molnár and A. Bousseksou, *Nat. Commun.*, 2013, 4, 2607.
- 17 M. Matsuda, H. Isozaki and H. Tajima, *Thin Solid Films*, 2008, **517**, 1465–1467.
- 18 C. Lochenie, K. Schötz, F. Panzer, H. Kurz, B. Maier, F. Puchtler, S. Agarwal, A. Köhler and B. Weber, J. Am. Chem. Soc., 2018, 140, 700–709.
- C. Bartual-Murgui, A. Akou, C. Thibault, G. Molnar, C. Vieu, L. Salmon and A. Bousseksou, *J. Mater. Chem. C*, 2015, 3, 1277–1285.
- 20 W.-Q. Gao, Y.-S. Meng, C.-H. Liu, Y. Pan, T. Liu and Y.-Y. Zhu, *Dalton Trans.*, 2019, **48**, 6323–6327.
- 21 J. Ru, F. Yu, P.-P. Shi, C.-Q. Jiao, C.-H. Li, R.-G. Xiong, T. Liu, M. Kurmoo and J.-L. Zuo, *Eur. J. Inorg. Chem.*, 2017, 2017, 3144–3149.

- K. E. Burrows, S. E. McGrath, R. Kulmaczewski,
  O. Cespedes, S. A. Barrett and M. A. Halcrow, *Chem. Eur. J.*, 2017, 23, 9067–9075.
- 23 I. A. Gass, S. Tewary, G. Rajaraman, M. Asadi, D. W. Lupton, B. Moubaraki, G. Chastanet, J.-F. Létard and K. S. Murray, *Inorg. Chem.*, 2014, 53, 5055–5066.
- A. Absmeier, M. Bartel, C. Carbonera, G. N. L. Jameson,
  F. Werner, M. Reissner, A. Caneschi, J. F. Letard and
  W. Linert, *Eur. J. Inorg. Chem.*, 2007, 3047–3054, DOI: 10.1002/ejic.200601096..
- 25 M. Muttenthaler, M. Bartel, P. Weinberger, G. Hilscher and W. Linert, *J. Mol. Struct.*, 2005, **741**, 159–169.
- 26 M. Seifried, C. Knoll, G. Giester, J. M. Welch, D. Müller and P. Weinberger, *Eur. J. Org. Chem.*, 2017, 2017, 2416– 2424.
- 27 M. Seifried, C. Knoll, G. Giester, M. Reissner, D. Müller and P. Weinberger, *Magnetochemistry*, 2016, 2, 1–13.
- 28 M. Nihei, Y. Yanai, I. J. Hsu, Y. Sekine and H. Oshio, Angew. Chem., Int. Ed., 2017, 56, 591–594.
- 29 C. Lochenie, J. Heinz, W. Milius and B. Weber, *Dalton Trans.*, 2015, 44, 18065–18077.
- 30 W. Bauer, C. Lochenie and B. Weber, *Dalton Trans.*, 2014, 43, 1990–1999.

- 31 A. Absmeier, M. Bartel, C. Carbonera, G. N. L. Jameson,
  P. Weinberger, A. Caneschi, K. Mereiter, J. F. Letard and
  W. Linert, *Chem. Eur. J.*, 2006, **12**, 2235–2243.
- 32 P. L. Franke and W. L. Groeneveld, *Transition Met. Chem.*, 1981, 6, 54–56.
- 33 T. Kamiya and Y. Saito, US3767667A, 1971.
- 34 T. Kamiya and Y. Saito, DE2147023A1, 1971.
- 35 O. H. Jústiz, R. Fernández-Lafuente, J. M. Guisán, P. Negri, G. Pagani, M. Pregnolato and M. Terreni, *J. Org. Chem.*, 1997, 62, 9099–9106.
- 36 D. Müller, C. Knoll, M. Seifried and P. Weinberger, Vib. Spectrosc., 2016, 86, 198–205.
- 37 P. Gütlich, A. Hauser and H. Spiering, Angew. Chem., Int. Ed. Engl., 1994, 33, 2024–2054.
- 38 A. Abhervé, M. J. Recio-Carretero, M. López-Jordà, J. M. Clemente-Juan, J. Canet-Ferrer, A. Cantarero, M. Clemente-León and E. Coronado, *Inorg. Chem.*, 2016, 55, 9361–9367.
- 39 D. D. Perrin and W. L. F. Armarego, *Purification of Laboratory Chemicals, 4th ed.*, Butterworth-Heinemann, Oxford, 1997.
- 40 I. Bruker, Bruker AXS Inc., Madison, Wisconsin, USA, 2012.