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Fine tuning of intra-lattice electron transfers through site doping in tetraoxolene-bridged iron honeycomb layers[†]

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The precise control of intra-lattice multiple electron transfers was demonstrated in the solvated and desolvated species of the tetraoxolene-bridged Fe honeycomb layer system, $(NPr_4)_2[Fe_2(Cl_2An)_3]$. (solv) ($Cl_2An^{n-} = 2,5$ -dichloro-3,6-dihydroxy-1,4-benzoquinonate; $NPr_4^+ =$ tetrapropylammonium cation), by the site-doping of the Cl_2An^{n-} bridging unit using X_2An^{n-} units with X = Br or F.

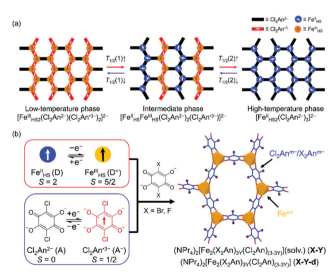
Exploring new stimuli-controllable functional systems is important in the field of materials science.^{1–4} These systems can be obtained by achieving thermally driven electron transfers (TDETs) in a material,⁵ which can successfully induce a state (or phase) change in materials at a transition temperature ($T_{1/2}$). Moreover, modifications of the physical properties of materials such as electronic conductivity,⁶ magnetic properties,⁷ elasticity,⁸ and the dielectric property can be attained.⁹ Currently, the relevant systems in the charge-transfer metal complexes, which are composed of electron-donor (D) and -acceptor (A) moieties,^{9–20} and organic charge transfer systems,^{21–24} have been investigated. Nevertheless, the systematic fine tuning of the variable charge states is still challenging; however, it can potentially be used for the application of these systems in molecular devices and sensors.^{1–3}

Recently, we have reported two kinds of tetraoxolenebridged iron honeycomb layered compounds, $(NPr_4)_2[Fe_2 (Cl_2An)_3] \cdot 2(acetone) \cdot H_2O$ (**0**; $Cl_2An^{n-} = 2,5$ -dichloro-3,6dihydroxy-1,4-benzoquinonate; $NPr_4^+ =$ tetrapropylammonium cation)¹⁵ and their desolvated compound $(NPr_4)_2[Fe_2(Cl_2An)_3]$ (**0-d**).¹⁶ The former exhibited a TDET at $T_{1/2a} = 236$ K between the phases of $[Fe^{III}_2(Cl_2An^{2-})(Cl_2An^{\cdot 3-})_2]^{2-}$ at $T < T_{1/2a}$ and $[Fe^{II}Fe^{II}(Cl_2An^{2-})_2(Cl_2-An^{\bullet 3-})]^{2-}$ at $T > T_{1/2a}$,¹⁵ while the latter underwent a stepwise TDET at $T_{1/2b} = 317$ K and $T_{1/2c} = 354$ K which changed phases from $[Fe^{III}_2(Cl_2An^{2-})(Cl_2An^{\bullet 3-})_2]^{2-}$ at $T < T_{1/2b}$ to $[Fe^{II}_2(Cl_2An^{2-})_3]^{2-}$ at $T > T_{1/2c}$ via the intermediate state of $[Fe^{II}Fe^{III}(Cl_2An^{2-})_2(Cl_2An^{\bullet 3-})]^{2-}$ at $T_{1/2b} < T < T_{1/2c}$ (Scheme 1a).¹⁶ In addition, compounds **0** and **0-d** are reversible via the solvation and desolvation processes, respectively. Hence, the systematic and continuous tuning of these phases were favorable due to being unprecedented in multiple TDET systems.

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Herein, we report a precise tuning of the multiple transition temperatures $T_{1/2a}$, $T_{1/2b}$, and $T_{1/2c}$ based on a chemical technique of site-doping for the charge-transfer materials **0** and **0-d**. A rational incorporation of the redox metal ions and bridging



Scheme 1 Schematic representations of the investigated tetraoxolenebridged Fe honeycomb layers. (a) The variations of charge-ordered states in **0** and **0-d**, where for **0**, $T_{1/2}(1)$ was $T_{1/2a}$ (*i.e.*, there was no hightemperature phase for **0**), and for **0-d**, $T_{1/2}(1)$ and $T_{1/2}(2)$ correspond to $T_{1/2b}$ and $T_{1/2c}$, respectively. (b) Schematic route for the syntheses of sitedoped solid solution compounds in this work.

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ligands into the charge-transfer skeleton is one of the efficient strategies in achieving the fine tuning of TDET.^{17,18,25-28} In this work, the Cl₂An^{*m*-} unit in **0** and **0-d** was replaced by X₂An^{*m*-} with X = F or Br in several ratios in the isostructural series where F₂An^{*m*-} = 2,5-difluoro-3,6-dihydroxy-1,4-benzoquinonate and Br₂An^{*m*-} = 2,5-dibromo-3,6-dihydroxy-1,4-benzoquinonate. Because the F₂An²⁻ and Br₂An²⁻ units act as stronger and weaker electron acceptors than Cl₂An²⁻, respectively, their solid-solution series revealed inverse trends in the doping rate dependence of $T_{1/2}$ s between the F-doped and Br-doped series, that is, proportional and inversely-proportional, respectively, despite being linearly variable in both cases.

The solvated compound series of the solid solutions $(NPr_4)_2[Fe_2(Cl_2An)_{(3-3Y)}(X_2An)_{3Y}]\cdot 2(acetone)\cdot H_2O$ (X-Y) as X = Br with Y = 0.10 (Br-0.10), 0.18 (Br-0.18), 0.35 (Br-0.35), 0.65 (Br-0.65) and X = F with Y = 0.06 (F-0.06), 0.12 (F-0.12), 0.17 (F-0.17), 0.24 (F-0.24), 0.31 (F-0.31), 0.42 (F-0.42) were synthesized using an identical procedure for 0. Specifically, H_2Br_2An or H_2F_2An was mixed into an H_2Cl_2An solution in various ratios (Scheme 1b),¹⁵ where the Y value was obtained from the halogen elemental analyses for the X-Y and their desolvated series, X-Y-d (ESI†). The Y doping ratio was approximately identical to the mixing ratio in the synthetic procedures.

All compounds of the solvated form (X-Y) were isostructural to **0** in the Y-doped region of $0.65 \ge Y$ and $0.42 \ge Y$ for X = Br and F, respectively. The compounds crystalized in the monoclinic $P2_1/c$ space group with an asymmetric unit containing two crystallographically independent Fe^{n+} ions, three $Cl_2An^{m-}/$ X_2An^{m-} ligands, and two NPr₄⁺ cations (Fig. S1 and Table S1, S2, ESI[†]). The Fe ions and Cl₂An/X₂An ligands formed a honeycomb anionic layer $[Fe_2(Cl_2An/X_2An)_3]^{2-}$ in the (100) plane. The occupancy ratio (*i.e.*, the X/Cl ratios) in the Cl₂An/X₂An ligands was fixed using the ratio obtained using the halogen elemental analyses (ESI[†]). The polycrystalline samples of the X-Y series were macroscopically evaluated using the powder X-ray diffractograms (PXRD) recorded at room temperature and compared with the simulated patterns determined using the single crystal XRD (SC-XRD) analyses (Fig. 1). We investigated the highly doped compounds beyond the doping rate of 0.65 < Y and 0.42 < Y for X = Br and F, respectively. The polycrystalline samples for the highly doped compounds and X₂An^{m-}-pure compounds had different PXRD patterns from that of 0. Unfortunately, the SC-XRD analyses for the highly doped samples did not succeed because of their low quality single crystals.

The desolvated series **X-Y-d** were obtained by using the samples of **X-Y**; $(NPr_4)_2[Fe_2(Cl_2An)_{(3-3Y)}(X_2An)_{3Y}]$ (X = Br; **Br-0.10-d** for *Y* = 0.10, **Br-0.18-d** for *Y* = 0.18, **Br-0.35-d** for *Y* = 0.35, **Br-0.65-d** for *Y* = 0.65, X = F; **F-0.06-d** for *Y* = 0.06, **F-0.12-d** for *Y* = 0.12, **F-0.17-d** for *Y* = 0.17, **F-0.24-d** for *Y* = 0.24, **F-0.31-d** for *Y* = 0.42) (Scheme 1b, Fig. S2, ESI†). The SC-XRD analysis at 103 K revealed that the **X-Y-d** compounds successfully changed to an isostructural 2D honeycomb layer structure with **0-d** (Fig. S3, S4 and Tables S3, S4, ESI†). Both the **Br-Y-d** and **F-Y-d** were crystallized in the monoclinic space group, $P2_1/n$, with an asymmetric unit containing one

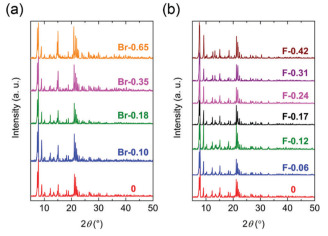


Fig. 1 PXRD patterns of $Br\mathchar`-Y$ (a) and $F\mathchar`-Y$ (b) measured at room temperature.

crystallographically unique Fe^{n+} (Fe \times 1), one and a half $\text{Cl}_2\text{An}^{m-}/\text{X}_2\text{An}^{m-}$ ligands, and one NPr_4^+ cation (Fig. S3, ESI†) NPr_4^+ was located between the honeycomb anionic layers $[\text{Fe}_2(\text{Cl}_2\text{An}/\text{X}_2\text{An})_3]^{2-}$ formed by the Fe ions and $\text{Cl}_2\text{An}^{m-}/\text{X}_2\text{An}^{m-}$ ligands, which were parallel to the (10–1) plane (Fig. S3, ESI†).

The oxidation states of the Fe^{*n*⁺} and X₂An^{*m*-} ligands for the series of **X-Y** and **X-Y-d** at 103 K were estimated from the local bond lengths of Fe–O and C–O in the Cl₂An^{*m*-}/X₂An^{*m*-} because their bond distances characteristically changed depending on their oxidation states.^{15,16,19,29} At 103 K, the electronic states of all the site-doped compounds were estimated to be a low-temperature state, which was similarly observed in **0** and **0-d** as ([(Fe^{III})₂(Cl₂An²⁻)(Cl₂An^{•3-})₂]²⁻) (see the ESI[†]).

The temperature dependence of the magnetic susceptibilities ($\chi = M/H$) of **Br-Y** (Y = 0.10, 0.18, 0.35, 0.65) and **F-Y** (Y = 0.06, 0.12, 0.17, 0.24, 0.31, 0.42) was measured by applying a 1 kOe dc field (H_{dc}) in the temperature range of 300 to 1.8 K (Fig. 2a and 3a and Fig. S5 and S6, ESI[†]). The $\chi_m T$ values at 300 K were in the range of 9.72-10.07 cm³ K mol⁻¹ and 9.89-11.28 cm³ K mol⁻¹ for the series of **Br-Y** and **F-Y**, respectively, which were varied in a narrow range (Table S5, ESI[†]). However, it is difficult to conclude that the variation of $\gamma_m T$ values at 300 K was caused by the electronic state modulations in the series because of the presence of a strong antiferromagnetic spin coupling between the Fe^{III} with S = 5/2 and $X_2An^{\bullet 3-}$ with S = 1/2 (Fig. S5, S6 and Table S5, ESI[†]).^{15,16,29–32} As the original **0** compound showed abrupt increases of χ_m and $\chi_m T$ at $T_{1/2a}$ = 236 K upon cooling concomitant with the one-step transition from the intermediate phase of [Fe^{II}Fe^{III}(Cl₂An²⁻)₂(Cl₂An^{•3-})]²⁻ to a low temperature phase of $[Fe^{III}_2(Cl_2An^{2-})(Cl_2An^{6-3-})_2]^{2-}$, all doped X-Y compounds exhibited similar transition features in $\chi_{\rm m}$ and $\chi_{\rm m}T$. However, their $T_{1/2a}$ shifted to lower and higher temperatures with increasing doping rate Y for Br-Y and F-Y, respectively, compared with $T_{1/2a}$ = 236 K for **0** (Fig. 2a and 3a); the $T_{1/2a}$ values reached 184 K (Δ 51 K from $T_{1/2a}$ in **0**) and 270 K (Δ 34 K) for **Br-0.65** and **F-0.31**, respectively. Specifically, $T_{1/2a}$ s in both Br-Y and F-Y linearly varied whereas the doping rate of

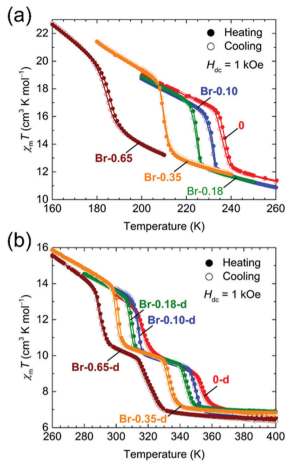


Fig. 2 Temperature dependence of $\chi_m T$ for **Br-Y** (a) and **Br-Y-d** (b) measured at $H_{dc} = 1 \text{ kOe} (\chi_m = M_m/H_{dc})$, where the narrow temperature ranges of 160–260 K and 260–400 K, respectively, were used.

Y had inversely proportional and proportional trends on Y, respectively (Fig. 4 and Fig. S7, ESI^{\dagger}), where the $T_{1/2a}$ temperatures were determined from the peak in the $d\chi_m T/dT vs. T$ plots and their error bars were estimated from the half-value width of the peaks (Fig. S8, ESI[†]). The slopes of the $T_{1/2a}$ variations were -77.3 and 110.3 K Y⁻¹ for **Br-Y** and **F-Y**, respectively. The **F-Y** series tended to have larger error bars with increasing Y than that of the Br-Y series, and $T_{1/2a}$ was not assigned in F-0.42, which could be due to the large random domain effect of F-X. Nevertheless, these trends of $T_{1/2a}$ in **Br-Y** and **F-Y** could be due to the site-doping of X_2An^{m-} with different electron acceptabilities, that is, $F_2An^{2-} > Cl_2An^{2-} > Br_2An^{2-}$ (electronegativity of halogen atom: F > Cl > Br).^{33,34} The magnetic profile in the entire temperature range from 300 to 1.8 K was similar to that for **0** excluding $T_{1/2a}$; however, the magnetic correlation lengths at low temperatures were slightly different from each other. Br-Y had a tendency to have a longer correlation length in highly doped compounds, whereas it was approximately inversed for F-Y (Fig. S5 and S6, ESI[†]). Furthermore, the superparamagnetic behavior of single-chain magnets was observed at lower temperatures as shown in the original compound 0 (Fig. S9, ESI⁺), which can be attributed to the ferrimagnetic chain motif of $[-Fe^{III}-(Cl_2An^{\bullet 3-}/X_2An^{\bullet 3-})-]$ (S_{Fe} = 5/2, S_{rad} = 1/2) separated by

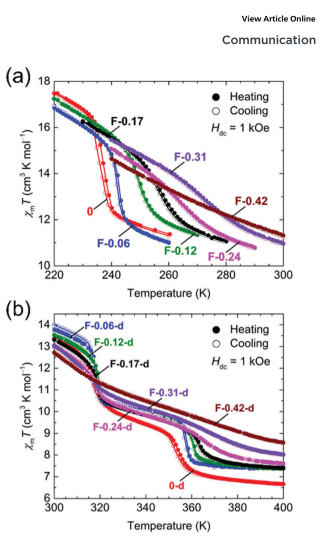


Fig. 3 Temperature dependence of $\chi_m T$ for **F-Y** (a) and **F-Y-d** (b) measured at $H_{dc} = 1 \text{ kOe} (\chi_m = M_m/H_{dc})$, where narrow temperature ranges of 220–300 K and 300–400 K, respectively, were used.

the diamagnetic Cl_2An^{2-}/X_2An^{2-} bridges in the honeycomb layer (low-temperature phase in Scheme 1a).¹⁵

The temperature dependence of the magnetic susceptibility for the doped series of the desolvated compounds X-Y-d was measured in the temperature range of 400–1.8 K. As the χ_m -and $\chi_m T - T$ plots for **0-d** showed a two-step transition (abrupt increasing of χ_m and $\chi_m T$ upon cooling) at $T_{1/2b}$ = 316 K and $T_{1/2c}$ = 353 K, corresponding to the variation of the electronic state from $[\text{Fe}_{2}^{\text{II}}(\text{Cl}_{2}\text{An}^{2-})_{3}]^{2-}$ at $T > T_{1/2c}$ to $[\text{Fe}^{\text{III}}_{2}(\text{Cl}_{2}\text{An}^{2-})(\text{Cl}_{2}\text{An}^{\bullet 3-})_{2}]^{2-}$ at $T < T_{1/2b}$ via an intermediate state of $[Fe^{II}Fe^{III}(Cl_2An^{2-})_2(Cl_2An^{\bullet 3-})]^{2-}$ at $T_{1/2b} < T < T_{1/2c}$ (Scheme 1a),¹⁶ the site-doped **X-Y-d** series showed similar twostep transitions. The error bar was larger in the highly doped species in F-X-d (Fig. 2b and 3b), owing to the characteristic domain effect. Moreover, the transition temperatures of $T_{1/2b}$ and $T_{1/2c}$ in **Br-Y-d** clearly shifted linearly to lower temperatures. On the other hand, the tendency in F-Y-d was slightly different from that of **F-Y**. The transition temperature of $T_{1/2c}$ in F-Y-d tended to shift to higher temperatures in the solid solution species with lower doping rates; however, $T_{1/2b}$ was approximately constant in the series. As mentioned above, the domain effect was characteristic in the series of F-Y-d and F-Y.

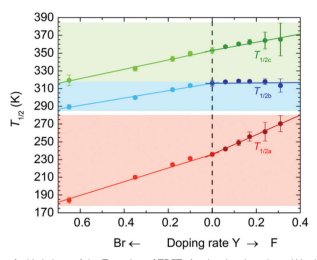


Fig. 4 Variations of the $T_{1/2}$ value of TDETs for the site-doped rate Y in the solid solution series of X-Y and X-Y-d, whereas the basis of Y = 0 corresponded to 0 and 0-d. The left and right scales represent **Br-Y/Br-Y-d** and **F-Y/F-Y-d**, respectively. The solid lines are least-square linear fitting lines for the respective series.

Hence, it is difficult to conclude that $T_{1/2b}$ was not affected by the site-doping of F_2An^{m-} . Although there was no clear reason, the transition temperatures in **F-Y-d** were slightly affected by the doping of the site-doping of F_2An^{m-} . The slope of the $T_{1/2b}$ and $T_{1/2c}$ variations *versus Y* was -41.7 and -52.7 K Y⁻¹ for **Br-Y-d** and 2.0 and 46.5 K Y⁻¹ and **F-Y-d** (Fig. S7, ESI⁺). The difference in tendencies can be explained by the electronacceptability of $F_2An^{2-} > Cl_2An^{2-} > Br_2An^{2-}$.

The site-doping of F_2An^{m-} influenced the magnetic phase of the solid-solution materials of **F-Y-d**. The **F-Y-d** species with $Y \le 0.24$ have a maximum of χ_m at approximately 23 K followed by a steep decrease after further cooling. This was indicative of an antiferromagnetic ground state as similarly found in **0-d** and **Br-Y-d** (Fig. S5, ESI†); however, the decrease of χ_m did not occur in the cases with higher doping rate. Instead, the superparamagnetic behavior of single-chain magnets was observed in the low-temperature phase of **F-Y-d** with 0.24 < *Y* (Scheme 1a).

In summary, we succeeded in the precise tuning of multiple TDET at three $T_{1/2}$ s transition temperatures through the chemical-site doping of two types of tetraoxolene-bridged Febased honeycomb layer compounds. This provided an "ondemand TDET" in a temperature range of 184 to 365 K.

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Conflicts of interest

There are no conflicts to declare.

Notes and references

- 1 O. Sato, Nat. Chem., 2016, 8, 644-656.
- 2 M. E. Guillermo and E. Coronado, Chem. Soc. Rev., 2018, 47, 533-557.
- 3 A. Dei, D. Gatteschi, C. Sangregorio and L. Sorace, Acc. Chem. Res., 2004, 37, 827–835.
- 4 E. Coronado, Nat. Rev. Mater., 2019, 5, 87-104.
- 5 J. S. Miller and K. S. Min, Angew. Chem., Int. Ed., 2009, 48, 262-272.
- 6 M. Mitsumi, T. Nishitani, S. Yamasaki, N. Shimada, Y. Komatsu, K. Toriumi, Y. Kitagawa, M. Okumura, Y. Miyazaki, N. Córska, A. Inaba, A. Kanda and N. Hanasaki, *J. Am. Chem. Soc.*, 2014, **136**, 7026–7037.
- 7 M. Nihei, Y. Sekine, N. Suganami, K. Nakazawa, A. Nakano, H. Nakano, Y. Murakami and H. Oshio, J. Am. Chem. Soc., 2011, 133, 3592–3600.
- 8 K. Boukheddaden, E. D. Loutete-Dangui, E. Codjovi, M. Castro, J. A. Rodriguéz-Velamazán, S. Ohkoshi, H. Tokoro, M. Koubaa, Y. Abid and F. Varret, *J. Appl. Phys.*, 2011, **109**, 013520.
- 9 N. Hoshino, F. Iijima, G. N. Newton, N. Yoshida, T. Shiga, H. Nojiri, A. Nakao, R. Kumai, Y. Murakami and H. Oshio, *Nat. Chem.*, 2012, 4, 921–926.
- 10 R. M. Buchanan and C. G. Pierpont, J. Am. Chem. Soc., 1980, 102, 4951-4957.
- 11 D. F. Li, R. Clérac, O. Roubeau, E. Harté, C. Mathonière, R. L. Bris and S. M. Holmes, *J. Am. Chem. Soc.*, 2008, **130**, 252–258.
- 12 D. Kiriya, H. C. Chang and S. Kitagawa, J. Am. Chem. Soc., 2008, 130, 5515–5522.
- 13 H. Miyasaka, N. Motokawa, T. Chiyo, M. Takemura, M. Yamashita, H. Sagayama and T. Arima, J. Am. Chem. Soc., 2011, 133, 5338–5345.
- 14 M. Nihei, Y. Okamoto, Y. Sekine, N. Hoshino, T. Shiga, I. P. C. Liu and H. Oshio, *Angew. Chem., Int. Ed.*, 2012, **51**, 6361–6364.
- 15 J. Chen, Y. Sekine, Y. Komatsumaru, S. Hayami and H. Miyasaka, Angew. Chem., Int. Ed., 2018, 57, 12043–12047.
- 16 J. Chen, Y. Sekine, A. Okazawa, H. Sato, W. Kosaka and H. Miyasaka, *Chem. Sci.*, 2020, **11**, 3610–3618.
- 17 K. Nakabayashi and H. Miyasaka, Chem. Eur. J., 2014, 20, 5121-5131.
- 18 K. Nakabayashi, M. Nishio and H. Miyasaka, *Inorg. Chem.*, 2016, 55, 2473–2480.
- 19 J. A. DeGayner, K. Y. Wang and T. D. Harris, J. Am. Chem. Soc., 2018, 140, 6550–6553.
- 20 W. Kosaka, Y. Takahashi, M. Nishio, K. Narushima, H. Fukunaga and H. Miyasaka, *Adv. Sci.*, 2018, 5, 1700526.
- 21 J. B. Torrance, A. Girlando, J. J. Mayerle, J. I. Crowley, V. Y. Lee and P. Batail, *Phys. Rev. Lett.*, 1981, **47**, 1747–1750.
- 22 S. Horiuchi, R. Kumai, Y. Okimoto and Y. Tokura, *Chem. Phys.*, 2006, 325, 78–91.
- 23 S. Horiuchi, R. Kumai, Y. Okimoto and Y. Tokura, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1999, 59, 11267–11275.
- 24 G. Saito and Y. Yoshida, Bull. Chem. Soc. Jpn., 2007, 80, 1-137.
- 25 M.-H. Zeng, B. Wang, X.-Y. Wang, W.-X. Zhang, X.-M. Chen and S. Gao, *Inorg. Chem.*, 2006, 45, 7069–7076.
- 26 M. Nishio, N. Hoshino, W. Kosaka, T. Akutagawa and H. Miyasaka, J. Am. Chem. Soc., 2013, 135, 17715–17718.
- 27 S. Chorazy, J. J. Stanek, W. Nogas, A. M. Majcher, M. Rams, M. Kozieł, E. Juszyńska-Gałązka, K. Nakabayashi, S.-I. Ohkoshi, B. Sieklucka and R. Podgajny, *J. Am. Chem. Soc.*, 2016, **138**, 1635–1646.
- 28 L. Liu, L. Li, M. E. Ziebel and T. D. Harris, J. Am. Chem. Soc., 2020, 142, 4705–4713.
- 29 J. A. DeGayner, I. R. Jeon, L. Sun, M. Dincă and T. D. Harris, J. Am. Chem. Soc., 2017, 139, 4175–4184.
- 30 A. Dei, D. Gatteschi, L. Pardi and U. Russo, *Inorg. Chem.*, 1991, 30, 2589–2594.
- 31 I. R. Jeon, B. Negru, R. P. Van Duyne and T. D. Harris, J. Am. Chem. Soc., 2015, 137, 15699–15702.
- 32 S. A. Sahadevan, A. Abhervé, N. Monni, D. P. Sáenz, J. R. Galán-Mascarós, J. C. Waerenborgh, B. J. C. Vieira, P. Auban-Senzier, S. Pillet, E. Bendeif, P. Alemany, E. Canadell, M. L. Mercuri and N. Avarvari, *J. Am. Chem. Soc.*, 2018, **140**, 12611–12621.
- 33 M. Atzori, S. Benmansour, G. Mínguez Espallargas, M. Clemente-León, A. Abhervé, P. Gómez-Claramunt, E. Coronado, F. Artizzu, E. Sessini, P. Deplano, A. Serpe, M. L. Mercuri and C. J. Gómez-García, *Inorg. Chem.*, 2013, 52, 10031–10040.
- 34 R. Murase, B. F. Abrahams, D. M. D'Alessandro, C. G. Davies, T. A. Hudson, G. N. L. Jameson, B. Moubaraki, K. S. Murray, R. Robson and A. L. Sutton, *Inorg. Chem.*, 2017, 56, 9025–9035.