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Electron transfer induced magnetic ordering of metal-cyanide magnets

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## Electron transfer induced magnetic ordering of metal-cyanide magnets<sup>†</sup>

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After over three decades of intense research effort, molecular cyanide magnets offer the prospect of low-temperature solution growth and stimuli-dependent magnetic properties. Central to the magnetic properties of cyanide compounds, electron transfer and spin coupling of structural cyanide networks are crucial because they control the magnetic ordering characteristics. However, the candidates of magnetically ordered molecules with controlled electron transfer are still very rare. Here we report atomic coordination induced control of electron transfer in solution grown metal-cyanide magnets, where the 2D FeSe layered material can react with the cyanide-chain dependent electron-accepting molecules, 7,7,8,8-tetracyanoquinodimethane and tetracyanoethylene. We demonstrate the electron transfer in Fe:cyanide networks, revealing its striking role in the magnetization and magnetic ordering. This study opens up a new avenue to design coordination magnets with high-temperature magnetic properties.

### Introduction

The cooperative magnetism of metal-cyanide molecular magnets arises from the localized spin exchange interaction between metal cations and cyanide radical anions,<sup>1</sup> which can be dictated through the control of the cyanide ligand dimension and metal-cyanide stoichiometry ratios.<sup>2–4</sup> Therefore, the electron transfer and molecular coupling can be designed to control magnetic hardening of molecular magnets for high Curie temperature.<sup>5,6</sup> The molecular magnet,  $V(TCNE)_x$  (TCNE = tetracyanoethylene) with ferrimagnetic ordering temperature ( $T_c$ ) beyond 400 K,<sup>7</sup> is the first metal-cyanide magnet ordered above room temperature. The abundant organometallic coordination structures render

metal-TCNE with metamagnetic, ferro- and photo-electric features to achieve the fundamental spin and charge coupling.<sup>8–12</sup> In this context, Fe(TCNE)-based magnets,  $Fe(TCNE)_2$  with  $T_c$  above 100 K, have been grown by the reaction between  $FeI_2$ ,  $Fe(CO)_5$  or  $FeCl_2(NCMe)_2$  and organic oxidizing agent TCNE.<sup>13–15</sup> However, it is challenging to control Fe and TCNE stoichiometry and electron transfer, which play an important role in the spin coupling of Fe-TCNE magnets to determine their magnetic ordering temperature.<sup>5</sup>

The mean field model is suggested to understand the magnetic properties in the insulating magnets, where the ordering temperature  $T_c$  could be enhanced by increasing the spin exchange interaction  $J$ , number of magnetic neighbors and Curie constants.<sup>16</sup> In a ferrimagnetic Fe and cyanide system,  $Fe(TCNX)_y$ , (TCNX is TCNQ or TCNE, TCNQ = 7,7,8,8-tetracyanoquinodimethane),<sup>17</sup> two kinds of molecules show similar coordination with Fe cations. The TCNE molecule has a short distance of two cyano radicals that could increase the spin exchange interaction as a bridge in  $Fe(TCNX)_y$ . Therefore, spin exchange interaction determined by atomic coordination is a critical factor in improving the  $T_c$  of  $Fe(TCNX)_y$  molecular magnets. Here, we report a solution reaction between 2D FeSe layered material and electron accepting molecules, TCNQ and TCNE, for the understanding of the electron transfer role in the magnetic ordering of metal-cyanide molecular magnets. The substitution reactions of 2D FeSe proceed through electron transfer from Fe to cyanide ligands, leading to  $Fe(TCNE)_y$  magnets with high Curie temperature, in contrast to  $M-(TCNE)_2$  magnets by chemical reactions using metal salts.<sup>14,18</sup> Magnetic characterization of these compounds reveals the significance of the stoichiometry and electron transfer in magnetic ordering of such molecular magnets. The magnetic order and electron transfer are influenced by the length of the bridge and stoichiometry ratios between the radical cyanide ligands (TCNQ or TCNE) and  $Fe^{(II)}$  cations. Through magnetic susceptibility and valence state analysis, the charge distribution in the Fe-TCNE network is taken into consideration for the mechanistic understanding of electron transfer induced magnetic ordering.

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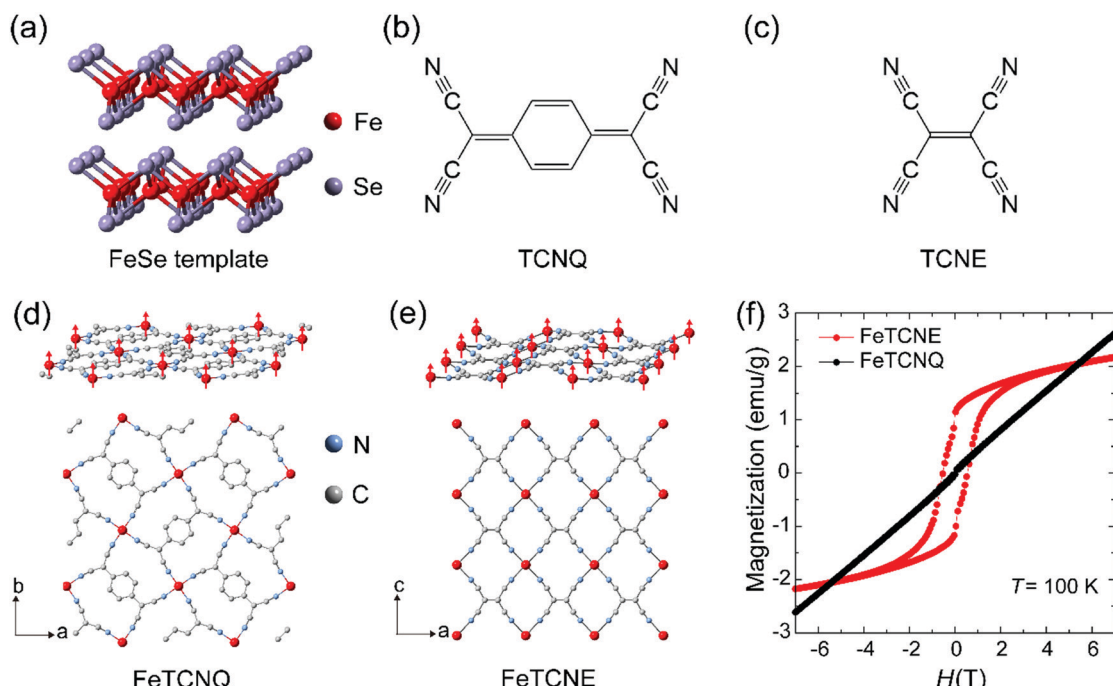


## Results

Fig. 1 shows the scheme to control ferrimagnetic exchange coupling order in molecular magnets *via* the reaction of 2D FeSe and molecular oxidizing agent TCNX (TCNQ and TCNE) cyanide ligands, which exhibit different ligand dimensions. A two-dimensional FeSe (Fig. 1a) layer serves as the iron-based precursor to chemically synthesize molecular Fe(TCNX)<sub>y</sub> magnets where magnetic order is originated from the spin coupling between Fe(II) cations and TCNX ligands. We reported the strong electronegativity of  $\text{C}\equiv\text{N}$  in TCNQ molecules for the synthesis of Fe-TCNQ molecular magnets (Fig. 1b). The TCNE molecule has the same four  $\text{C}\equiv\text{N}$  radicals as TCNQ while a lack of an aromatic carbon ring makes TCNE a shorter chain dimension (Fig. 1c). As an electron acceptor with strong electronegativity, TCNE performs well in the procedure of synthesizing Fe-TCNE molecular magnets by substituting Se in FeSe. The TCNX ligand accepts one electron to form a radical TCNX<sup>-</sup> oxidation state with an unpaired spin ( $S = 1/2$ ), distributing over the whole molecule.<sup>19</sup> In Fe(TCNX)<sub>y</sub> molecular magnets, one Fe atom loses two electrons and forms Fe<sup>2+</sup> cations with four unpaired spins ( $S = 2$ ), where the Fe<sup>2+</sup> cation coordinates with four  $\text{C}\equiv\text{N}$  radicals in Fe-TCNQ<sup>20,21</sup> (Fig. 1d) and six  $\text{C}\equiv\text{N}$  radicals in Fe-TCNE<sup>18,22,23</sup> (Fig. 1e). In the layer of Fe-TCNX, the antiferromagnetic spin coupling between the Fe<sup>2+</sup> cation ( $S = 2$ ) and TCNX<sup>-</sup> ( $S = 1/2$ ) presents a ferrimagnetic order due to their

unbalanced opposing spin numbers. Fig. 1d and e demonstrate the in-plane crystal structure and spin order of the Fe<sup>2+</sup> cation where the spin distributions of TCNX are not shown. The Fe(TCNE)<sub>y</sub> magnet possesses a compact Fe spin network due to its smaller cyanide ligand compared to Fe(TCNQ)<sub>y</sub>. Under the ligand design rule, Fe(TCNE)<sub>y</sub> should possess a higher ferrimagnetic ordering temperature due to its shorter ligand connection among spins between Fe<sup>2+</sup> cations using TCNE<sup>-</sup>.<sup>13</sup> The magnetization hysteresis ( $MH$ ) loops at 100 K for Fe(TCNE)<sub>y</sub> feature ferrimagnetic characteristics (Fig. 1f), which is drastically different from that of the Fe(TCNQ)<sub>y</sub> system. The paramagnetism of Fe(TCNQ)<sub>y</sub> at 100 K is consistent with its Curie temperature of 60 K. Because of its relatively short TCNE ligands, larger orbital overlap of unpaired electrons enhances the ferrimagnetic coupling between Fe<sup>2+</sup> cations and TCNE<sup>-</sup> radicals. Thus, Fe(TCNE)<sub>y</sub> is selected for further studies.

The electron transfer from Fe to TCNE proceeds *via* the formation of Fe(II) ions and reduced TCNE radicals in a stoichiometry ratio, which leads to the tunability on spin coupling and magnetization of Fe:TCNE controlled under solution-based reaction conditions. The reaction of FeSe and TCNE is thermodynamically driven, while a higher temperature involves an enhanced Fe:TCNE stoichiometry ratio  $y$  (Table S1, ESI<sup>†</sup>) for enhanced ferrimagnetic coupling. Fig. 2a compares the  $MH$  data of Fe(TCNE)<sub>y</sub> samples synthesized at 368 K and 410 K. For the sample synthesized at 410 K, the  $MH$  loop features a wide magnetic hysteresis with a coercivity of 7500 Oe at 50 K.



**Fig. 1** Crystal structures and magnetism of Fe-TCNX (X = Q and E). (a) The FeSe template shows the two-dimensional layered structure where four Se atoms surround Fe as the tetrahedral coordination in each layer. (b and c) Molecular structures of TCNQ and TCNE demonstrate that the molecular length of TCNE (4.189 Å) is smaller than that of TCNQ (7.757 Å). (d and e) Single-layer structures of Fe-TCNX illustrate the in-plane coordination of the Fe atom with cyano radicals. The bindings between layers and solvent molecules are not shown for clarity. Since molecules serve as bridges in the Fe network, all spins on Fe orient ferromagnetically in the ground state. Due to the smaller molecule size, the ferromagnetic coupling in Fe-TCNE with closer exchange interaction is stronger than that in Fe-TCNQ. (f) Magnetic field dependent magnetism of Fe-TCNE shows the large ferromagnetic hysteresis at 100 K above the Curie temperature of Fe-TCNQ of 60 K.



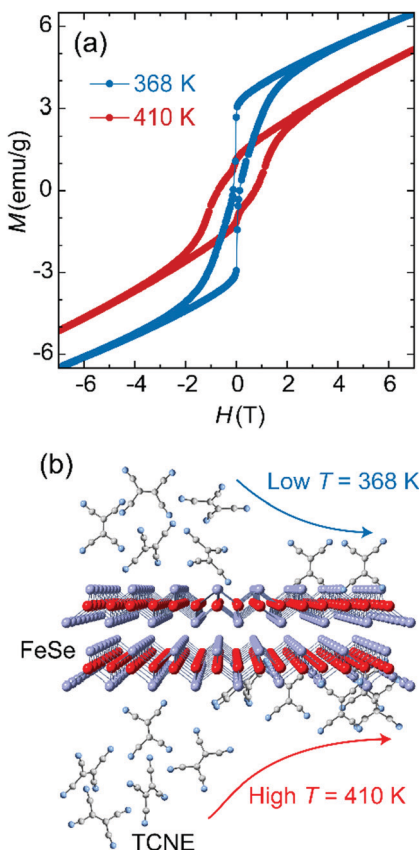


Fig. 2 Temperature effects on synthesis of molecular ferromagnet  $\text{Fe}(\text{TCNE})_y$ . (a) Magnetic hysteresis loops of  $\text{Fe}(\text{TCNE})_y$  at 50 K show that higher synthesis temperature improves the coercivity and homogeneity. (b) Reaction progress of FeSe and TCNE could be activated by heat energy. High enough synthesis temperature can improve the substitution reaction to obtain ferrimagnetic  $\text{Fe}(\text{TCNE})_y$ .

When increasing the magnetic field to 3 T, the magnetic hysteresis behavior fades away and the magnetism tends to be paramagnetic. However, once the synthesis temperature decreases to 368 K, the  $MH$  hysteresis loop narrows down, in which coercivity reduces to 1200 Oe and shows a kink at around zero magnetic field, in comparison to the sample synthesized at 410 K (Fig. 2a). Therefore, a high reaction temperature proceeds toward the product formation by building up  $\text{Fe}(\text{TCNE})_y$  coordination (Table S1, ESI<sup>†</sup>).

In addition, a low reaction temperature leads to a low coercivity that originates from a weak spin coupling interaction. The thermal energy improves the ferrimagnetism by enhancing the exchange interaction *via* more TCNE ligand connections with Fe atoms. Energy-dispersive X-ray spectroscopy analysis (EDS) on the  $\text{Fe}(\text{TCNE})_y$  sample synthesized at 368 K results in the TCNE stoichiometry of 1.00 (Table S1, ESI<sup>†</sup>). However, for a sample synthesized at 410 K, the  $\text{Fe}(\text{TCNE})_y$  molecular structure possesses a higher TCNE stoichiometry of 3.75 (Table S1, ESI<sup>†</sup>) that connects among  $\text{Fe}(\text{II})$  cations to build up a strong ferrimagnetic coupling. The structure change by the TCNE stoichiometry can also be seen from the X-ray diffraction patterns (Fig. 3 and Fig. S1, ESI<sup>†</sup>). A wide range of TCNE concentrations in

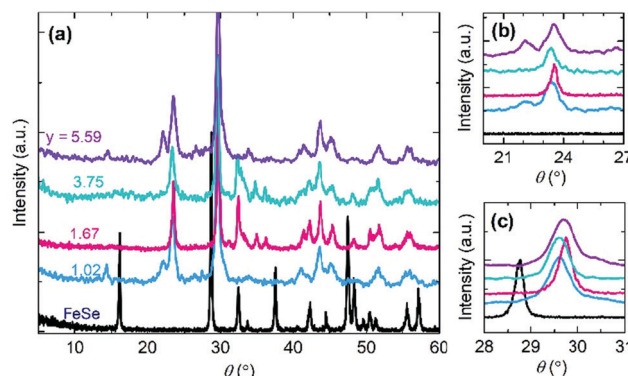


Fig. 3 X-Ray diffraction patterns of  $\text{Fe}(\text{TCNE})_y$  synthesized at 410 K by a FeSe template. (a) XRD patterns of the whole range from 5° to 60° present the structure evolution from FeSe to  $\text{Fe}(\text{TCNE})_y$  ( $y = 1.02, 1.67, 3.75$  and 5.59). (b) Enlarged area of XRD patterns from 20° to 27°. (c) Enlarged area of XRD patterns from 28° to 31°.

$\text{Fe}(\text{TCNE})_y$  samples may be attributed to the multiple valence states of TCNE and the mixed stoichiometries under different reaction conditions. The IR spectra of  $\text{Fe}(\text{TCNE})_y$  present multiple absorption peaks at the  $\nu_{\text{CN}}$  region (Fig. S2, ESI<sup>†</sup>). Particularly for the TCNE-rich sample, extra broad absorption at  $2329 \text{ cm}^{-1}$  indicates that neutral TCNE molecules<sup>24</sup> exist in the  $\text{Fe}(\text{TCNE})_y$  sample. The thermal energy plays a vital role in the substitution reaction for the formation of  $\text{Fe}(\text{TCNE})_y$ . Fig. 2b schematically illustrates the dynamic process of the substitution reaction by TCNE ligands in FeSe. High reaction temperature promotes the reaction for the formation of the  $\text{Fe}(\text{TCNE})_y$  structure, while a sluggish reaction kinetics occurs at low temperature. A lower reaction temperature leads to uncompleted and few substitutions in FeSe that may form the magnetization kink at zero field.

The electron transfer between Fe and TCNE ligands appears to be an important role in the formation of molecular magnets. The other effective approach to control the electron transfer is to define an appropriate amount of TCNE acceptor in the reaction process. Through controlling the TCNE reactant amount in the reaction solution, the TCNE ratio  $y$  in  $\text{Fe}(\text{TCNE})_y$  varies in a large range from 1.02 to 5.59. Fig. 3 shows the X-ray diffraction (XRD) patterns of  $\text{Fe}(\text{TCNE})_y$  synthesized at 410 K with a variable TCNE ratio  $y$ . After the substitution reaction, the XRD patterns have considerable changes that a lot of new peaks appear compared to that of the FeSe template (Fig. 3a). At a lower angle, a small peak arises in the  $\text{Fe}(\text{TCNE})_{1.02}$  compound indicating a larger crystal lattice than that of FeSe. Meanwhile, most peaks in the XRD patterns of  $\text{Fe}(\text{TCNE})_y$  cannot be tracked from the XRD pattern of FeSe, which could be attributed to the dramatic structure change. Two peaks appear at around 24° in the XRD patterns of  $\text{Fe}(\text{TCNE})_y$  where there should be no peaks for FeSe (Fig. 3b). In addition, the peak of highest intensity in  $\text{Fe}(\text{TCNE})_y$  at around 30° locates at a higher angle than that in FeSe (Fig. 3c). No apparent peak shift law was observed for these  $\text{Fe}(\text{TCNE})_y$  compounds that may result from the inhomogeneity of substitution. The complicated evolution of the XRD patterns of  $\text{Fe}(\text{TCNE})_y$  implies the significant crystal structure change that influences the spin coupling interaction.

The complex magnetic properties of the  $\text{Fe}(\text{TCNE})_y$  products at 100 K (Fig. 4a) reveal the effect of  $\text{Fe:TCNE}$  stoichiometry ratio  $y$  on electron transfer. For a low TCNE stoichiometry ( $y = 1.02$ ), the  $MH$  loop is almost linear and it shows no magnetic hysteresis, suggesting paramagnetic nature. When  $y = 1.67$  and 3.75, as we discussed in Fig. 2, the  $\text{Fe}(\text{TCNE})_y$  network shows a large magnetic hysteresis and remnant magnetization. By further improving the TCNE stoichiometry, the  $MH$  loop is linear without hysteresis. The stoichiometry significance can also be observed by the temperature dependent susceptibility (MT) of the as-synthesized  $\text{Fe}(\text{TCNE})_y$  (Fig. 4b). It shows the magnetic susceptibilities of four samples with different TCNE stoichiometry ratios. For the two  $\text{Fe}(\text{TCNE})_y$  samples where the paramagnetic phase is dominated (Fig. 4a), the  $M-T$  data show much lower susceptibilities compared to those of ferrimagnetic samples (Fig. S5 and S6, ESI<sup>†</sup>). Ferrimagnetic  $\text{Fe}(\text{TCNE})_y$  displays a high susceptibility at around 150 K. A small susceptibility upturn at low temperature could be attributed to a possible paramagnetic phase like Se. The tunability of ferromagnetism in  $\text{Fe}(\text{TCNE})_y$  is realized through the adjustment of the quantity of TCNE in the synthesis to engineer FeSe templates at different levels. When the reaction temperature remains constant, more TCNE ligands

promote the formation of the  $\text{Fe}(\text{TCNE})_y$  network from a FeSe template. There is a rationale for the critical role of TCNE stoichiometry in the magnetism of Fe-TCNE, where the Se atoms are gradually and partially substituted by TCNE ligands. In order to understand the different reaction stages of FeTCNE, here we propose a magnetic cluster model to picture the magnetic properties after a substitution reaction under different TCNE concentrations. At a low concentration of TCNE, the reaction does not occur uniformly to transform FeSe into  $\text{Fe}(\text{TCNE})_y$  assembly. In this case, magnetic clusters of  $\text{Fe}(\text{TCNE})_y$  form short-range ferrimagnetic order and present weak magnetic behavior (stage I, Fig. 4c). This situation satisfies the sample of  $y = 1.02$  in Fig. 4a that is paramagnetic in nature with nonlinear magnetization at around zero field. The XRD of the sample of  $y = 1.02$  (Fig. 3a) indicates the lattice expansion due to the incidence of TCNE molecules. For the other two samples showing ferrimagnetism, the stoichiometry of the TCNE ligands enables the long-range ferrimagnetic order due to the uniform substitution (stage II and III, Fig. 4d and e). However, the sample of  $y = 3.75$  with highly substituted TCNE ligands results in a slightly disordered coordination among layers in  $\text{Fe}(\text{TCNE})_y$ , leading to the paramagnetic behavior (stage III, Fig. 4e). The XRD pattern of the sample of  $y = 3.75$  (Fig. 3) confirms its disordered feature in the Fe-TCNE network, which could further interrupt the spin lattice and break the ferrimagnetic order (stage IV, Fig. 4f). The strong charm of this work is not only to synthesize ferrimagnetic  $\text{Fe}(\text{TCNE})_y$  of high  $T_c$ , but also the various molecular assembly structures by TCNE substitution at different levels. In stage I, only a few TCNE ligands are coordinated in the FeSe framework that is analogous to a molecular heterostructure in different concentrations of TCNE. This hybrid structure may also be the reason for the kink behavior observed in the  $MH$  loops of  $\text{Fe}(\text{TCNE})_y$  (Fig. 2a and Fig. S3, S4, ESI<sup>†</sup>), especially in those samples synthesized at low temperature (Fig. S3, ESI<sup>†</sup>). Such molecular heterostructures can tune the magnetism by interface and lead to high temperature molecular magnets, which is often contributed to the optimum stoichiometry between TCNE radical anions and metal cations.<sup>5</sup>

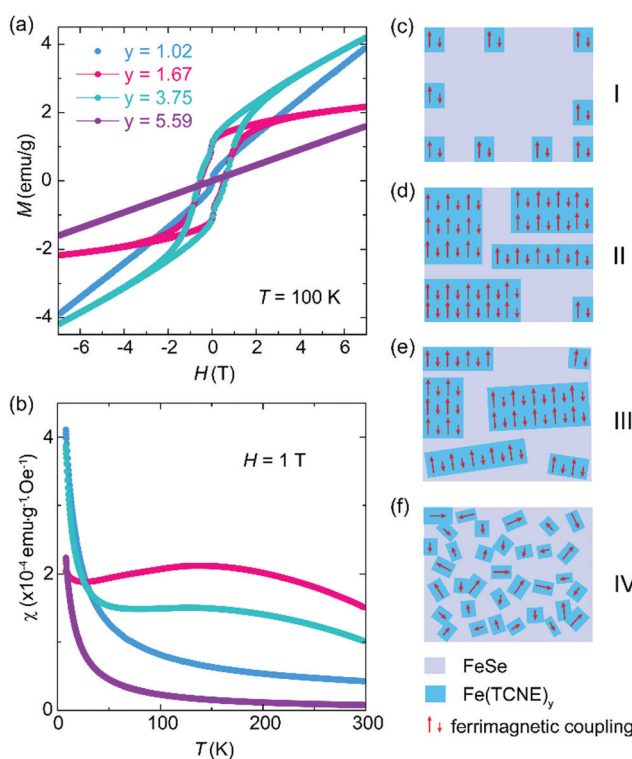


Fig. 4 The evolution of the magnetic hysteresis loop of  $\text{Fe}(\text{TCNE})_y$  samples synthesized at 410 K by different TCNE stoichiometry  $y = 1.02, 1.67, 3.75$  and 5.59 which correspond to four stages of spin configuration in  $\text{Fe}(\text{TCNE})_y$ . (a) Too low ( $y = 1.02$ ) or too high ( $y = 5.59$ ) TCNE stoichiometry makes a paramagnetic phase, and only appropriate ratios of  $y = 1.67$  and 3.75 result in a ferrimagnetic coupling. (b) The temperature dependent susceptibilities of these four samples in (a) show low (paramagnetic) and high (ferrimagnetic) susceptibilities. (c-f) Schematic figures for different reaction stages of diverse spin configurations according to the TCNE reactant concentration at the synthesis temperature of 410 K.

## Conclusions

In conclusion, we design and synthesize a high temperature Fe-cyanide-based molecular magnet *via* a chemical reaction between 2D FeSe and molecular oxidizing agent TCNX. The shorter TCNE molecules enhance the spin coupling between Fe and cyanide, presenting a higher ordering temperature. The electron transfer induced spin coupling in a molecular magnet enables the design of tunable magnetic interaction by the control of the spin lattice and stoichiometry ratios. The reaction conditions, such as temperature and TCNE concentrations, are vital in tuning the final structure of  $\text{Fe}(\text{TCNE})_y$  after the substitution reaction that determines the long-range ferrimagnetic coupling and ordering temperature of even possible above room temperature (Fig. S7, ESI<sup>†</sup>). The selection of the FeSe precursor complex and small cyanide molecules provides a new pathway to synthesize high temperature molecular magnets and molecular



interface devices that benefit the potential applications in spin electronics.<sup>25</sup>

## Experimental section

### Sample preparation

The synthesis of Fe-TCNE was conducted in a 20 ml glass bottle which contains 0.8 mmol FeSe powder, 0.4–6.4 mmol TCNE and 10 ml acetonitrile solvent. All chemicals were sealed in the glass bottle which later on was heated at a synthesis temperature of 368 K or 410 K on a hotplate. A small magnetic stir bar was added into the reaction solution and stirred at 1000 rpm to increase the reaction rate. Three days later, the product was washed with acetonitrile and then centrifuged to condense in the bottom of the centrifuge tube. After the product was dried in a fume hood, the following measurements were taken on those samples.

### Structural and morphologic characterization

X-Ray diffractions of powder samples were carried out on a Rigaku Ultima IV (40 kV, 44 mA) to characterize the crystal structure evolution due to the TCNE substitution. A Carl Zeiss AURIGA (200 kV) with an Oxford Energy-dispersive X-ray Spectrometer (EDS) was used to collect the surface morphology and element ratios of the samples obtained under different reaction conditions.

### Magnetic susceptibility and hysteresis loop measurements

Magnetic property measurements were conducted on a Physical Properties Measurement System (PPMS) equipped with a Vibrating Sample Magnetometer (VSM) option. After cooling at zero magnetic field, magnetic susceptibility under different magnetic fields was measured from 8 K to 300 K. Magnetic hysteresis loops were measured at fixed temperatures by sweeping the magnetic field from 7 T to 7 T.

### Infrared spectra

A Bruker Alpha IR spectrometer with ALPHA-P Platinum ATR module (diamond crystal) is used to obtain transmission ATR-FTIR spectra under an argon atmosphere in a glove box.

## Conflicts of interest

There are no conflicts to declare.

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