

MIT Open Access Articles

Triphenylene-Bridged Trinuclear Complexes of Cu: Models for Spin Interactions in Two-Dimensional Electrically Conductive Metal-Organic Frameworks

The MIT Faculty has made this article openly available. *Please share* how this access benefits you. Your story matters.

Citation: Yang, Luming et al. "Triphenylene-Bridged Trinuclear Complexes of Cu: Models for Spin Interactions in Two-Dimensional Electrically Conductive Metal-Organic Frameworks." Journal of the American Chemical Society 141, 26 (June 2019): 10475–10480 © 2019 American Chemical Society

As Published: http://dx.doi.org/10.1021/jacs.9b04822

Publisher: American Chemical Society (ACS)

Persistent URL: https://hdl.handle.net/1721.1/128206

Version: Author's final manuscript: final author's manuscript post peer review, without

publisher's formatting or copy editing

Terms of Use: Article is made available in accordance with the publisher's policy and may be subject to US copyright law. Please refer to the publisher's site for terms of use.



Triphenylene-Bridged Trinuclear Complexes of Cu: Models for Spin Interactions in Two-Dimensional Electrically Conductive MOFs

Luming Yang, Xin He, and Mircea Dincă*

Department of Chemistry, Massachusetts Institute of Technology, 77 Massachusetts Avenue, Cambridge, Massachusetts 02139, United States

Supporting Information Placeholder

ABSTRACT: Reaction of 2,3,6,7,10,11-hexahydroxytriphenylene (HHTP) and 2,3,6,7,10,11-hexaaminotriphenylene (HATP) with [Cu(Me₃tacn)]²⁺ (Me₃tacn = 1,4,7-trimethyl-1,4,7-triazacyclononane) produces trigonal tricopper complexes [(Me₃tacnCu)₃(HOTP)]³⁺ (1) and [(Me₃tacnCu)₃(HITP)]⁴⁺ (2) (HOTP, HITP = hexaoxy- and hexaimino-triphenylene, respectively). These trinuclear complexes are molecular models for spin exchange interactions in the two-dimensional conductive metal-organic frameworks (MOFs) copper hexaoxytriphenylene (Cu₃HOTP₂) and copper hexaiminotriphenylene (Cu₃HITP₂). Whereas complex 1 is isolated with HOTP³⁻ bearing the same oxidation state as found in the oxy-bridged MOF, the triply oxidized HITP³⁻ found in Cu₃HITP₂ is unstable with respect to disproportionation in the molecular model. Indeed, magnetic measurements reveal ligand-centered radical character for 1 and a closed-shell structure for 2, in agreement with the redox state of the ligands. All neighboring spins are antiferromagnetically coupled in 1 and 2. These results help probe metal-ligand-metal interactions in conductive MOFs and provide potential inspiration for the synthesis of other two-dimensional materials with delocalized electrons.

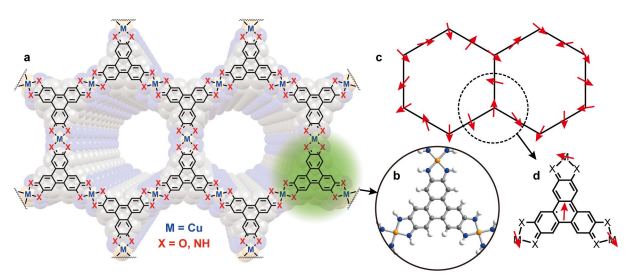


Figure 1. (a) Structural and chemical representation of typical 2D conductive MOFs, with depiction of the graphite-like honeycomb structure as well as HXTP (X = O, I) ligand-centered radical. The highlighted part illustrates the trinuclear metal – ligand monomeric unit modeled here as shown in (b). (c) Spin lattice of typical 2D conductive MOFs, with arrows showing randomized spin centers. (d) Spin structure of the monomeric unit depicted in (b), showing metal- and ligand-centered radicals.

INTRODUCTION

Multifunctional spintronic devices that respond to electrical, magnetic, and chemical stimuli have drawn much attention in recent years due to their important role in downsizing logic circuits.¹⁻³ One class of materials that show promise towards such applications are two-dimensional (2D) electrically conductive metal-organic frameworks (MOFs) made from paramagnetic ions that in principle

respond to electrical, magnetic, as well as chemical stimuli.⁴⁻
⁷ These 2D MOFs have layered graphene-like honeycomb structures, where the vertices in the hexagonal tiles are the tritopic organic ligands and the transition metals ions bridging the ligands reside on the edges of each hexagon (Figure 1).^{4,10,17} Because the ligands and metals in MOFs can be independently modified or chosen, these materials present opportunities that are not easily achieved with graphite or other 2D materials, allowing for control over porosity, conductivity, and spin state. However, the

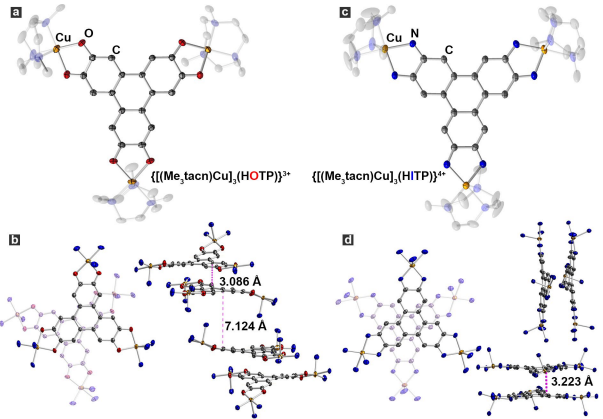


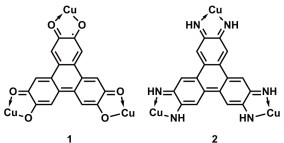
Figure 2. Crystal structures of 1 and 2, showing (a, c) single cationic fragments containing Cu₃HXTP motifs, (b, d) top (left) and side (right) views of the dimeric packing modes in unit cells. Thermal ellipsoids are plotted at 50% probability level for elements other than hydrogen. All anions and solvent molecules are omitted for clarity. The L backbones are also omitted in (b) and (d). The red dotted lines indicate selected inter-molecular C-C distances.

extended structures and compositional complexity also translate into complicated electronic structures that give rise to electrical and magnetic properties that are still poorly understood despite numerous experimental and computational studies. 8,9,18-20 One powerful means to interrogate the properties of solids has been dimensional reduction, wherein the independent synthesis and study of smaller building blocks provide insight into the electronic structure of the parent solids. 21,22 The 2D conductive materials described above make excellent targets for this strategy. Here, we report trinuclear Cu complexes bridged by the same trigonal ligands found in the MOFs and study their redox and magnetic properties.

RESULTS AND DISCUSSION

In a manner mimicking the isolation of small clusters related to Prussian blue analogs, 23 we employed Cu²+ complexes capped with the neutral multidentate κ_3 -chelating ligand 1,4,7-trimethyl-1,4,7-triazacyclononane (Me₃tacn) to target the MOF model complexes. Importantly, the typical coordination mode of Me₃tacn keeps the copper magnetic orbital (dx²-y²) in the aromatic plane of the ligand, as is also likely the case in Cu₃HXTP₂ (X = O, I). Because in the extended MOFs the ligand is formally triply oxidized – and thus bears a formal oxidation state of -3 after six-fold deprotonation of the amino or hydroxy groups – we targeted the same oxidation state in the molecular complexes (Scheme 1, left). In view of the expected rich redox reactivity

of [M₃HOTP]ⁿ⁺ complexes,²⁴⁻²⁶ the target compounds [(Me₃tacnCu)₃(HXTP)]ⁿ⁺ were synthesized under inert atmosphere followed by stoichiometric oxidations with three equivalent of oxidants. Thus, reaction of $[(Me_3tacn)Cu](BF_4)_2$ with 2,3,6,7,10,11hexahydroxytriphenylene (HHTP) under N2, followed by deprotonation and oxidation with three equivalents of ferrocenium tetrafluoroborate (FcBF₄) yielded $[(Me_3tacnCu)_3(HOTP)](BF_4)_3 (\mathbf{1}\cdot (BF_4)_3)$ as dark navy crystals in 73% yield (Scheme S1). Unexpectedly, a similar procedure 2,3,6,7,10,11-hexaaminotriphenylene instead of HHTP did not lead to the isolation of the desired [(Me3tacnCu)3(HITP)]3+ complex. Instead, upon reaction of HATP with [(Me3tacn)Cu]2+ and oxidation by three equivalents of FcBF4, an initially dark blue-green solution changes to a dark blue solution whereupon only $[(Me_3tacnCu)_3(HITP)](BF_4)_4$ (2·(BF₄)₄) could be isolated (Scheme 1, right; also Scheme S1). This more electrondeficient species containing the quadruply oxidized ligand HITP²⁻ suggests that [(Me₃tacnCu)₃(HITP)]³⁺ is unstable with respect to disproportionation, a redox behavior that diverges from that of Cu₃HITP₂ under otherwise similar synthetic conditions. The identities of 1 and 2 were confirmed by high-resolution electrospray ionization/mass spectrometry (ESI/MS) and microelemental analyses. The former gave m/z values of 1285.3 for 1 ([M+H]+, M = $[(C_9H_{21}N_3Cu)_3(C_{18}H_6O_6)](BF_4)_3)$ and m/z = 1365.4 for 2 $([M+H]^+, M = [(C_9H_{21}N_3Cu)_3(C_{18}H_{12}N_6)](BF_4)_4$, see also Supporting Information).



Scheme 1. Lewis structures of selected resonance forms of 1 and 2, illustrating the charge states and spin structures of HXTP (X = O, I), respectively. The capping Me_3 tacn ligands are omitted for clarity.

Single crystals of 1 and 2 suitable for X-ray diffraction (XRD) analysis were obtained by layering or diffusing vapors of t-butyl-methyl-ether to solutions of the respective complexes in 4:1 mixtures of dichloroethane:methanol at -5 $^{\circ}$ C. Compounds 1 and 2 crystallize in space groups $P\bar{1}$ and P21/c, respectively (Fig. 2a, c). In the solid state, both 1 and 2 pack in dimeric fashion, with closest C-C distances of 3.086(5) and 3.223(6) Å between the dimers (Fig. 2b, d; right). A top-down view of the dimers reveals staggered conformations for both complexes (Fig. 2b,d). In both 1 and 2, the longest Cu-N_{Me2}tacn bonds are the axial ones, as expected due to the Jahn-Teller distortion, and confirming the co-planarity of the Cu $d_{x^2-v^2}$ magnetic orbital with the HXTP plane. For metal-semiguinone complexes, the C-O bond length is often related to the degree of oxidation of the semiquinone fragment. Shorter C-O bonds indicate more oxidized (quinone-type) character whereas longer C-O bonds correspond to reduced (catechol-type) character.²⁷ For 1, the average C-O bond length is 1.283(4) Å, in very good agreement with the value of 1.288(2) Å reported for an established copper bis-semiguinonate complex.²⁸ In contrast, 2 exhibits an average C-N bond length of 1.305(6) Å, shorter than that of a structurally relevant bisiminosemiquinonate copper azophenine complex (1.327(3) Å).²⁹ Although subtle, the difference between the shorter average C-N bond length in 2 relative to the established iminosemiquinonate Cu complex is consistent with a higher degree of oxidation and partial iminoquinone character in 2.

Cyclic voltammetry (CV) provided further support for assigning formal oxidation states in 1 and 2 and offered clues into the degree of electron delocalization in the trinuclear complexes. CV experiments for complex 1 conducted in 0.1 M propylene carbonate solutions of tetrabutylammonium hexafluorophosphate (TBAPF₆) under N₂ reveal three reversible reduction events at -o.80 V, -o.45V, -o.09 V relative to the ferrocenium/ferrocene (Fc+/Fc) couple (Fig. 3a). All three values are in line with expected ligandcentered redox couples: HOTP^{3-/4-}, HOTP^{4-/5-}, and HOTP⁵⁻ ^{/6-}, the last corresponding to the formation of neutral [(Me₃tacnCu)₃(HOTP)].^{25,30} In the framework of the classical theory of mixed-valence compounds, which relates electrochemical redox potentials to the degree of electronic delocalization between mixed-valence centers,31,32 the potential difference between the HOTP5-/6- and HOTP3-/4couples, $\Delta E = 0.71$ V, equates to a comproportionation constant,33 Kc, of 1012.0. This large value indicates that the three redox centers - the three semiquinone/hydroquinone moieties of each HOTP ligand - are fully delocalized and form a Robin-Day class III mixed-valent compound,34 as may

be expected for three formal radicals, all centered on a single organic moiety. A similar analysis for 2 is made difficult by the irreversibility of the first reduction event, presumably the HITP^{2-/3-} couple, which occurs at -0.51 V (Fig. 3b). The irreversible reduction of 2 is in line with rapid disproportionation of the [(Me₃tacnCu)₃(HITP)]³⁺ species, which further prevents access to more reduced complexes. Although the strong degree of charge delocalization within HOTP is in-line with the extended delocalization observed in the respective HOTP-based MOF, inaccessible Cu-based redox events with either 1 or 2 prevent the assessment of charge delocalization between the *metal* centers, as mediated by HOTP or HITP.

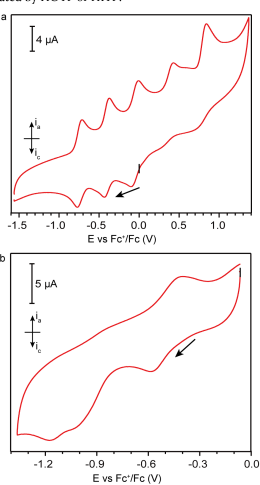


Figure 3. Cyclic voltammograms of 1 (a) and 2 (b) in 0.1 M TBAPF $_6$ – propylene carbonate under N_2 . Scanning directions are marked by arrows. Open circuit potentials are indicated by black tick marks.

Magnetometry provided critical information about the electronic communication between metals and/or ligands in complexes 1 and 2. Variable temperature direct current magnetic susceptibility measurements indicated antiferromagnetic coupling for both complexes, as revealed by rapid decreases of $\chi_M T$ below 50 K in an applied field of 2.5 kOe (Fig. 4a). For 1, a room temperature $\chi_M T$ value of 1.79 cm3mol-1K is close to the value expected for four independent $S = \frac{1}{2}$ spins (1.50 cm³mol⁻¹K, with q = 2.0), representing three non-interacting Cu^{2+} ions and one $S = \frac{1}{2}$ HOTP³⁻ organic radical. Because of a significant contribution from temperature independent paramagnetism (TIP) (see Fig. S2 and related discussion for assignment of TIP), the room-temperature $\chi_M T$ value for 2 is 2.32 cm³mol¬¹K, higher than expected for three uncoupled Cu²+ ions connected through the diamagnetic HITP²- (see Fig. S3 for $\chi_M T$ vs. T plot where the contributions of TIP were corrected for). The plateau $\chi_M T$ value for 2 of 0.42 cm³mol¬¹K at 1.8 K is nevertheless close to the value expected for a S=1/2 spin system (0.375 cm³mol¬¹K with g=2.0). This suggests that the ground spin state of 2 is S=1/2, as would be expected for three antiferromagnetically coupled Cu²+ ions at low temperature.

To assess the magnitude of the spin exchange interactions, the temperature-dependent $\chi_M T$ data were fitted with Heisenberg-Dirac-Van Vleck (HDVV) Hamiltonians, $\mathbf{H_1}$ for $\mathbf{1}$ and $\mathbf{H_2}$ for $\mathbf{2}$ (see also Fig. S4):

$$\begin{aligned} \mathbf{H}_1 &= -2J_1(\mathbf{S}_{\text{Cu}1}\mathbf{S}_{\text{L}} + \mathbf{S}_{\text{Cu}2}\mathbf{S}_{\text{L}}) - 2J_2\mathbf{S}_{\text{Cu}3}\mathbf{S}_{\text{L}} \\ \mathbf{H}_2 &= -2J_1(\mathbf{S}_{\text{Cu}1}\mathbf{S}_{\text{Cu}2} + \mathbf{S}_{\text{Cu}1}\mathbf{S}_{\text{Cu}3}) - 2J_2\mathbf{S}_{\text{Cu}2}\mathbf{S}_{\text{Cu}3} \end{aligned}$$

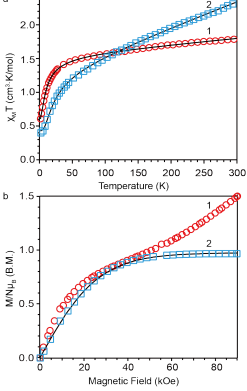


Figure 4. (a) Temperature-dependent $\chi_M T$ (H = 2.5 kOe) and (b) field-dependent magnetization (T = 1.8 K) curves for 1 and 2. The solid black lines are the best fits as described in the text.

Good fits were obtained for both complexes by using two J values representing only the spin interactions between nearest neighbors. For $\mathbf{1}$, fitting with contribution from TIP and intermolecular coupling, $\mathbf{z}J'$, as well as two g values for copper- and ligand-centered spins gives $J_1 = J_2 = -2.76$ cm⁻¹ (with $g_{\text{Cu}} = 2.06$, $g_{\text{L}} = 1.98$, $\chi_{\text{TIP}} = 8.86 \cdot 10^{-4}$ cm³ mol⁻¹, $\mathbf{z}J' = -0.309$ cm⁻¹) (Fig. 4a) (see Fig. S5 and related discussion on the choice of g). This indicates that the ligand-based radical is coupled to the three Cu²⁺ ions through weak antiferromagnetic interactions of similar strength. As such, the ground spin state of $\mathbf{1}$ is $S = \mathbf{1}$, with the trigonal symmetry preserved in the spin structure despite a geometric

distortion from perfect planarity. For 2, the best fit of the temperature-dependent $\chi_M T$ data gave $J_1 = J_2 = -6.99$ cm⁻¹, with $g_{\text{Cu}} = 2.12$ and $\chi_{TIP} = 3.71\cdot 10^{-3}$ cm³mol⁻¹ (Fig. 4a). To assess the possibility that paramagnetic impurities contribute to the magnetic properties of 1 and 2, we fitted the data by adding S = ½ Cu²⁺ impurities. This produced fits with similar parameters as for the pure samples above, with negligible contribution from the S = ½ impurities (Fig. S6 and related discussions). The lack of crystalline impurities is also supported by the powder X-ray diffraction analysis for 1 and 2 (see Fig. S7 and related discussions).

The ground spin states of complexes 1 and 2 were confirmed by variable field magnetization measurements (Fig. 4b). For 1, the magnetization does not saturate even at 90 kOe and continues to grow after reaching an inflection point of ~1 B.M. at approximately 40 kOe, a behavior that is typically associated with the presence of low-lying excited spin states.³⁵ The magnetization curve for 1 further deviates from typical Brillouin behavior, likely because of additional intermolecular antiferromagnetic interactions that lower the expected saturation value of 2 B.M. for an S = 1 system (see Fig. S8 and related discussion in the Supporting Information). In contrast, the magnetization of complex 2 follows typical Brillouin-type behavior and shows a saturation value of 1 B.M., as expected for an S = 1/2 system. Fitting of this data with a Brillouin function gives q = 2.0, a slightly smaller value than usually observed for Cu²⁺, again likely owing to weak intermolecular antiferromagnetic interactions. Low temperature electron paramagnetic resonance (EPR) provided further insight on the electronic structure of 1 and 2. For 1, an isotropic S = 1/2 signal at g = 1/22.00 and an axial S = 1/2 signal with g_{\parallel} = 2.25, g_{\perp} = 2.05, and hyperfine coupling to Cu^{2+} (I = 3/2) with $A_{\parallel} \sim 30$ G were observed (Fig. S9). This agrees with the observation of single unpaired spins on both HOTP and Cu²⁺ from magnetometry measurements in 1. For 2, only an axial S = 1/2 signal with g_{\parallel} = 2.25, g_{\perp} = 2.05 is observed, which gives a good fit with Cu²⁺ hyperfine coupling constants of A_{\parallel} = 40 G and A_{\perp} = 413 G, suggesting the presence of S = 1/2 spins coupled to Cu^{2+} in 2. This is in good agreement with the magnetometry data that suggests the absence of an HITP-based radical in 2.

Density functional theory (DFT) calculations carried out for the cationic fragments [(Me3tacnCu)3HOTP]3+ and [(Me₂tacnCu)₂HITP]⁴⁺ corroborate magnetometry data and provide further insight into the orbital interactions involved in the spin exchange mechanisms in 1 and 2. For both complexes, DFT calculations agree with the proposed spin density distribution derived from magnetometry data above. The magnetic orbitals of the copper centers have the same symmetry as the $d_{x^2-y^2}$ atomic orbitals, and lie in the plane of the molecule (Fig. 5). For both 1 and 2, the oxygen and nitrogen atoms of the HXTP (X = O, I) ligands have primarily the same sign of spin density as the copper atoms, suggesting ferromagnetic exchange due to orbital orthogonality. Meanwhile, the dominating antiferromagnetic interaction in both complexes is embodied through spin polarization or antiferromagnetic superexchange mechanisms in the p- π systems of the ligands.

Altogether, the structural, electrochemical, and magnetic data support a ligand-based $S = \frac{1}{2}$ radical in 1, which therefore represents a first example of a molecular complex modeling electronic interactions in a conductive MOF, in this case Cu_3HOTP_2 . Notably, even though other examples

of isolable M_3 HXTP-type complexes (X = O, N, S) exist, none of these show spin density on both the metal and the ligand. ^{24–26,30,32,36} Although 1 is a truncated model of an idealized single-sheet of Cu_3 HOTP₂, which to our knowledge has not been accessed yet, if the antiferromagnetic interactions observed in 1 are reproduced in a single sheet of Cu_3 HOTP₂, the latter may exhibit properties consistent with a conductive 2D ferrimagnet. ^{20,37} Verifying such rare behavior rests entirely with efforts to generate and exfoliate sufficiently large single crystals of Cu_3 HOTP₂, an area of ongoing interest in our laboratory.

In the same vein, it should be noted that **2** satisfies the classical geometric spin frustration criterion of $J_1 = J_2 < 0$ in a spin triangle, as proposed by Kahn.³⁸ Although it has been established that true geometric spin frustration cannot be achieved in molecular complexes due to fast geometric relaxation,^{39,40} the type of interactions observed in **2** bring about the possibility of a geometrically frustrated solid state material, where structural relaxation to lower symmetry is quenched by lattice rigidity. As with **1**, if it exists, spin frustration is likely to be observed only in single, charged sheets of extended MOFs of the type $M_3(HITP)_2$, with no inter-sheet interactions.^{41,42} Once again, the challenge in accessing such 2D structures rests squarely in the synthetic realm and highlights the importance of controlling crystal growth for 2D conductive MOFs.

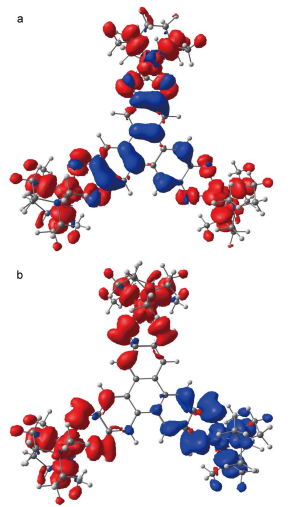


Figure 5. Calculated spin density isosurfaces (0.0007 a.u.) of (a) 1 and (b) 2. Red and blue represent spin-up and spin-down densities.

CONCLUSIONS

In conclusion, we present the synthesis, electrochemical, and magnetic characterization of two new trinuclear copper complexes that mimic the smallest building blocks of conductive 2D MOFs Cu₃HOTP₂ and Cu₃HITP₂. Whereas the oxygen-based complex retains a ligand-based radical similar to the corresponding Cu₃HOTP₂, the nitrogen-based ligand radical disproportionates, thereby diverging from the formal oxidation state present in Cu₃HITP₂. In both complexes, the metal and/or ligand-based electronic spins are coupled through weak antiferromagnetic interactions. These results portend potentially exciting long-range magnetic interactions in isolated single sheets of the respective 2D MOFs, substantiating the importance of future efforts to isolate such sheets.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website.

Details of experimental procedures, single crystal X-ray crystallography, fitting of magnetic data, and computational methods (PDF)

X-ray crystal structure of 1 (CIF)

X-ray crystal structure of 2 (CIF)

AUTHOR INFORMATION

Corresponding Author

*mdinca@mit.edu

Notes

The authors declare no competing financial interests.

ACKNOWLEDGMENT

This work was supported by the Army Research Office (grant number W911NF-17-1-0174). We thank Professor T. David Harris for fruitful discussions.

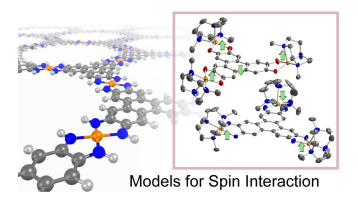
REFERENCES

- (1) Wolf, S. A.; Awschalom, D. D.; Buhrman, R. A.; Daughton, J. M.; von Molnár, S.; Roukes, M. L.; Chtchelkanova, A. Y.; Treger, D. M. Spintronics: A Spin-Based Electronics Vision for the Future. *Science* **2001**, *294*, 1488–1495.
- (2) Awschalom, D. D.; Flatté, M. E. Challenges for Semiconductor Spintronics. *Nat. Phys.* **2007**, *3*, 153–159.
- (3) Ballav, N.; Wäckerlin, C.; Siewert, D.; Oppeneer, P. M.; Jung, T. A. Emergence of On-Surface Magnetochemistry. *J. Phys. Chem. Lett.* **2013**, *4*, 2303–2311.
- (4) Hmadeh, M.; Lu, Z.; Liu, Z.; Gándara, F.; Furukawa, H.; Wan, S.; Augustyn, V.; Chang, R.; Liao, L.;

- Zhou, F.; Perre, E.; Ozolins, V.; Suenaga, K.; Duan, X., Dunn, B.; Yamamto, Y.; Terasaki, O.; Yaghi, O. New Porous Crystals of Extended Metal-Catecholates. *Chem. Mater.* **2012**, *24*, 3511–3513.
- (5) Kambe, T.; Sakamoto, R.; Hoshiko, K.; Takada, K.; Miyachi, M.; Ryu, J. H.; Sasaki, S.; Kim, J.; Nakazato, K.; Takata, M.; Nishihara, H.; l. π-Conjugated Nickel Bis(Dithiolene) Complex Nanosheet. *J. Am. Chem. Soc.* **2013**, *1*35, 2462–2465.
- (6) Huang, J.; He, Y.; Yao, M. S.; He, J.; Xu, G.; Zeller, M.; Xu, Z. A Semiconducting Gyroidal Metal-Sulfur Framework for Chemiresistive Sensing. *J. Mater. Chem. A* **2017**, *5*, 16139–16143.
- (7) Wäckerlin, C.; Chylarecka, D.; Kleibert, A.; Müller, K.; Iacovita, C.; Nolting, F.; Jung, T. A.; Ballav, N. Controlling Spins in Adsorbed Molecules by a Chemical Switch. *Nat. Commun.* **2010**, *1*, 1–7.
- (8) Chakravarty, C.; Mandal, B.; Sarkar, P. Bis(Dithioline)-Based Metal-Organic Frameworks with Superior Electronic and Magnetic Properties: Spin Frustration to Spintronics and Gas Sensing. *J. Phys. Chem. C* **2016**, 120, 28307–28319.
- (9) Mandal, B.; Sarkar, P. A New Two-Dimensional Metal-Organic Framework with High Spin-Filtering Efficiency. *Phys. Chem. Chem. Phys.* **2015**, 17, 17437–17444.
- (10) Sheberla, D.; Sun, L.; Blood-Forsythe, M. A.; Er, S.; Wade, C. R.; Brozek, C. K.; Aspuru-Guzik, A.; Dincă, M. High Electrical Conductivity in Ni3(2,3,6,7,10,11- Hexaiminotriphenylene)2, a Semiconducting Metal-Organic Graphene Analogue. *J. Am. Chem. Soc.* 2014, 136, 8859–8862.
- (11) Dou, J. H.; Sun, L.; Ge, Y.; Li, W.; Hendon, C. H.; Li, J.; Gul, S.; Yano, J.; Stach, E. A.; Dincă, M. Signature of Metallic Behavior in the Metal-Organic Frameworks M3(Hexaiminobenzene)2 (M = Ni, Cu). *J. Am. Chem. Soc.* 2017, 139, 13608–13611.
- (12) Sheberla, D.; Bachman, J. C.; Elias, J. S.; Sun, C. J.; Shao-Horn, Y.; Dincă, M. Conductive MOF Electrodes for Stable Supercapacitors with High Areal Capacitance. *Nat. Mater.* 2017, *16*, 220–224.
- (13) Feng, D.; Lei, T.; Lukatskaya, M. R.; Park, J.; Huang, Z.; Lee, M.; Shaw, L.; Chen, S.; Yakovenko, A. A.; Kulkarni, A.; Xiao, J.; Fredrickson, K.; Tok, J. B.; Zou, X.; Cui, Y.; Bao, Z.; Robust and Conductive Two-Dimensional Metal-Organic Frameworks with Exceptionally High Volumetric and Areal Capacitance. *Nat. Energy* 2018, 3, 30–36.
- (14) Benmansour, S.; Abhervé, A.; Gómez-Claramunt, P.; Vallés-García, C.; Gómez-García, C. J. Nanosheets of Two-Dimensional Magnetic and Conducting Fe(II)/Fe(III) Mixed-Valence Metal-Organic Frameworks. ACS Appl. Mater. Interfaces 2017, 9, 26210–26218.
- (15) Dong, R.; Han, P.; Arora, H.; Ballabio, M.; Karakus, M.; Zhang, Z.; Shekhar, C.; Adler, P.; Petkov, P. S.; Erbe, A.; Mannsfeld, S. C. B.; Felser, C.; Heine, T.; Bonn, M.; Feng, X.; Cánovas, E.; High-Mobility Band-like Charge Transport in a Semiconducting Two-Dimensional Metal-Organic Framework. *Nat. Mater.* 2018, 17, 1027–1032.
- (16) Zhang, X.; Vieru, V.; Feng, X.; Liu, J.; Zhang, Z.; Na, B.; Shi, W.; Wang, B.; Powell, A. K.; Chibotaru, L. F.; Gao, S.; Cheng, P.; Long, J. R.; Influence of Guest

- Exchange on the Magnetization Dynamics of Dilanthanide Single-Molecule-Magnet Nodes within a Metal-Organic Framework. *Angew. Chem., Int. Ed.* **2015**, 54, 9861–9865.
- (17) Campbell, M. G.; Sheberla, D.; Liu, S. F.; Swager, T. M.; Dincă, M. Cu3(Hexaiminotriphenylene)2: An Electrically Conductive 2D Metal-Organic Framework for Chemiresistive Sensing. *Angew. Chem., Int. Ed.* 2015, 54, 4349–4352.
- (18) Zhao, M.; Wang, A.; Zhang, X. Half-Metallicity of a Kagome Spin Lattice: The Case of a Manganese Bis-Dithiolene Monolayer. *Nanoscale* **2013**, 5, 10404–10408.
- (19) Zhou, Q.; Wang, J.; Chwee, T. S.; Wu, G.; Wang, X.; Ye, Q.; Xu, J.; Yang, S. W. Topological Insulators Based on 2D Shape-Persistent Organic Ligand Complexes. *Nanoscale* **2015**, *7*, 727–735.
- (20) Li, W.; Sun, L.; Qi, J.; Jarillo-Herrero, P.; Dincă, M.;
 Li, J. High Temperature Ferromagnetism in π Conjugated Two-Dimensional Metal-Organic
 Frameworks. *Chem. Sci.* 2017, 8, 2859–2867.
- (21) Long, J. R.; Williamson, A. S.; Holm, R. H. Dimensional Reduction of Re6Se8Cl2: Sheets, Chains, and Discrete Clusters Composed of Chloride-Terminated [Re6Q8]2+(Q = S, Se) Cores. *Angew. Chem., Int. Ed.* 1995, 34, 226–229.
- (22) Tulsky, E. G.; Long, J. R. Dimensional Reduction: A Practical Formalism for Manipulating Solid Structures. Chem. Mater. 2001, 13, 1149–1166.
- (23) Heinrich, J. L.; Berseth, P. A.; Long, J. R. Molecular Prussian Blue Analogues: Synthesis and Structure of Cubic Cr4Co4(CN)12 and Co8(CN)12 Clusters. *Chem. Commun.* 1998, 4, 1231–1232.
- (24) Grange, C. S.; Meijer, A. J. H. M.; Ward, M. D. Trinuclear Ruthenium Dioxolene Complexes Based on the Bridging Ligand Hexahydroxytriphenylene: Electrochemistry, Spectroscopy, and near-Infrared Electrochromic Behaviour Associated with a Reversible Seven-Membered Redox Chain. *Dalt. Trans.* 2010, 39, 200–211.
- (25) Suenaga, Y.; Inada, H.; Inomata, M.; Yamaguchi, R.; Okubo, T.; Maekawa, M.; Kuroda-Sowa, T. Crystal Structure and Characterization of Trinuclear Cobalt(III) Complex with 2,3,6,7,10,11-Hexahydroxytriphenylene. *Chem. Lett.* 2014, 43, 562–564.
- (26) Hoshino, N.; Akutagawa, T. A Trinuclear Iron(III) Complex of a Triple Non-Innocent Ligand toward Spin-Structured Molecular Conductors. *Chem. - A Eur. J.* 2018, No. Iii.
- (27) Pierpont, C. G.; Lange, C. W. The Chemistry of Transition Metal Complexes Containing Catechol and Semiquinone Ligands; 2007; Vol. 41.
- (28) Ovcharenko, V. I.; Gorelik, E. V.; Fokin, S. V.; Romanenko, G. V.; Ikorskii, V. N.; Krashilina, A. V.; Cherkasov, V. K.; Abakumov, G. A. Ligand Effects on the Ferro- to Antiferromagnetic Exchange Ratio in Bis(o-Semiquinonato)Copper(II). *J. Am. Chem. Soc.* 2007, 120, 10512–10521.
- (29) Schweinfurth, D.; Khusniyarov, M. M.; Bubrin, D.; Hohloch, S.; Su, C.-Y.; Sarkar, B. Tuning Spin–Spin Coupling in Quinonoid-Bridged Dicopper(II) Complexes through Rational Bridge Variation. *Inorg. Chem.* 2013, 52, 10332–10339.

- (30) Barthram, A. M.; Reeves, Z. R.; Jeffery, J. C.; Ward, M. D. Polynuclear Osmium-Dioxolene Complexes: Comparison of Electrochemical and Spectroelectrochemical Properties with Those of Their Ruthenium Analogues. J. Chem. Soc. Dalt. Trans. 2000, No. 18, 3162–3169.
- (31) Richardson, D. E.; Taube, H. Mixed-Valence Molecules: Electronic Delocalization and Stabilization. Coord. Chem. Rev. 1984, 60, 107–129.
- (32) Sakamoto, R.; Kambe, T.; Tsukada, S.; Takada, K.; Hoshiko, K.; Kitagawa, Y.; Okumura, M.; Nishihara, H. Π-Conjugated Trinuclear Group-9 Metalladithiolenes With a Triphenylene Backbone. *Inorg. Chem.* **2013**, 52, 7411–7416.
- (33) Sutton, J. E.; Taube, H. Metal to Metal Interactions in Weakly Coupled Mixed-Valence Complexes Based on Ruthenium Ammines. *Inorg. Chem.* **1981**, 20, 3125–3134.
- (34) Robin, M. B.; Day, P. Mixed Valence Chemistry-A Survey and Classification; 1968; pp 247–422.
- (35) Menage, S.; Vitols, S. E.; Bergerat, P.; Codjovi, E.; Kahn, O.; Girerd, J. J.; Guillot, M.; Solans, X.; Calvet, T. Structure of the Linear Trinuclear Complex Hexakis(Acetato)Bis(2,2'-Bipyridine)Trimanganese (II). Determination of the J Electron-Exchange Parameter through Magnetic Susceptibility and High-Field Magnetization Measurements. *Inorg. Chem.* 1991, 30, 2666–2671.
- (36) Kambe, T.; Tsukada, S.; Sakamoto, R.; Nishihara, H. Expanding Family of π-Conjugated Trinuclear Dithiolenes: The Cases of Group 8 (RuII) and 10 (NiII and PtII) Metals. *Inorg. Chem.* **2011**, 50, 6856–6858.
- (37) Liu, J.; Sun, Q. Enhanced Ferromagnetism in a Mn₃C₁₂N₁₂H₁₂ Sheet. *ChemPhysChem* **2015**, *16*, 614–620.
- (38) Kahn, O. *Molecular Magnetism*; VCH Publishers, Inc., 1993.
- (39) Tsukerblat, B. S.; Kuyavskaya, B. Y.; Belinskii, M. I.; Ablov, A. V.; Novotortsev, V. M.; Kalinnikov, V. T. Antisymmetric Exchange in the Trinuclear Clusters of Copper (II). *Theor. Chim. Acta* **1975**, *38*, 131–138.
- (40) Cage, B.; Cotton, F. A.; Dalal, N. S.; Hillard, E. a.; Rakvin, B.; Ramsey, C. M. Observation of Symmetry Lowering and Electron Localization in the Doublet-States of a Spin-Frustrated Equilateral Triangular Lattice: Cu₃(O₂C₁6H₂3)·1.2C6H₁2. *J. Am. Chem. Soc.* 2003, 125, 5270–5271.
- (41) Hagemann, I. S.; Huang, Q.; Gao, X. P. A.; Ramirez, A. P.; Cava, R. J. Geometric Magnetic Frustration in Ba₂Sn₂Ga₃ZnCr₇O₂₂: A Two-Dimensional Spinel Based Kagomé Lattice. *Phys. Rev. Lett.* **2001**, *86*, 894–897.
- (42) Shores, M. P.; Bartlett, B. M.; Nocera, D. G. Spin-Frustrated Organic-Inorganic Hybrids of Lindgrenite. *J. Am. Chem. Soc.* **2005**, *127*, 17986–17987.



For Table of Contents Only